

REPORT DRXTH-TE-CR-82179

SURFACE SAMPLING TECHNIQUES

Bruce E. Goodwin James R. Aronson Robert P. O'Neil Margaret A. Randel Emmett M. Smith

ARTHUR D. LITTLE, INC. CAMBRIDGE, MA 02140

SEPTEMBER, 1982

FINAL REPORT Volume I Distribution Unlimited Cleared for Public Release



prepared for

U.S. Army Toxic and Hazardous Materials Agency, Aberdeen Proving Ground, Maryland 21010

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7. Authorrei Bruce E. Goodwin		6. CONTRACT OR GRANT NUMBER(*)
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SUMMARY

Under Contract No. DAAK11-81-C-0014 to the U. S. Army Toxic and Hazardous Materials Agency, Arthur D. Little, Inc. has developed sampling protocols for the determination of explosives/explosive residues on building materials surfaces. The Army's need for this sampling and analysis capability has arisen in connection with the release of surplus government property (e.g., former ammunition plants) and the specified requirement that as part of these release programs, a determination be made as to whether the property can be released for unrestricted use. In the case of buildings known or suspected to have been contaminated with explosives, the Army is seeking for this purpose sampling and analytical procedures which would permit: (1) rapid qualitative determination with 90% confidence of the presence/absence of the compounds of interest down to a level of 5 $\mu g/10~cm^2$ in a given building, and (2) if any of these compounds is detected, precise, accurate quantitative determination of the amount of contamination down to the same 5 $\mu g/10~cm^2$ level.

This study resulted in the development of several methods for the sampling and analysis of explosives/explosive residues on building materials surfaces. A method for qualitative determination based on detection of charge-transfer complexes formed between the explosive/explosive residue and a visualization reagent applied to the surface has been evaluated in the field at two Army Ammunition Plants. A method for quantitative determination based on solvent extraction of samples collected in the field followed by high pressure liquid chromatographic analysis has been evaluated on samples prepared in the laboratory, and found to give promising results. The theoretical and practical feasibility of another method for qualitative determination based on UV irradiation of a suspected contaminated surface with subsequent detection of any explosives/explosive residues present has been demonstrated. This approach may provide the means for "scanning" an area to determine whether explosives are present on a real-time basis. Further development of this approach is recommended.

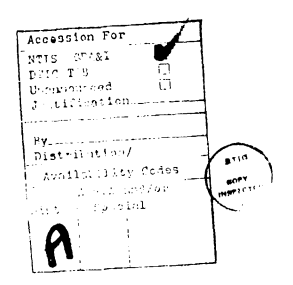


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I. INTRODUCTION

Under Contract No. DAAK11-81-C-0014 to the U.S. Army Toxic and Hazardous Materials Agency, Arthur D. Little, Inc. has developed sampling protocols for the determination of explosives/explosive residues on building materials surfaces. The Army's need for this sampling and analysis capability has arisen in connection with the release of surplus government property (e.g., former ammunition plants) and the specified requirement that as part of these release programs, a determination be made as to whether the property can be released for unrestricted use. In the case of buildings known or suspected to have been contaminated with explosives, the Army is seeking for this purpose sampling and analytical procedures which would permit: (1) rapid qualitative determination with 90% confidence of the presence/ absence of the compounds of interest down to a level of 5 μ g/10 cm² in a given building, and (2) if any of these compounds is detected, precise, accurate quantitative determination of the amount of contamination down to the same 5μ g/10 cm² level.

The specific compounds and surface types of interest were the following:

Compounds

- 1. 2,4,6-trinitrotoluene (TNT)
- 2. 2,4- and 2,6-dinitrotoluene
- 3. Cyclotrimethylenetrinitramine (RDX)
- 4. Pentaerythrite tetranitrate (PENT)
- 5. Nitroglycerine
- 6. 2,4,6-trinitrophynylmethylnitramine (Tetryl)
- 7. Diphenylamine
- 8. 1,3,5-trinitrobenzene
- 9. 2,4-dinitrophenol
- 10. Cd
- 11. Pb
- 12. Hg
- 13. Cr + 3
- 14. Cr+6

Surface Types

- 1. Concrete -- unpainted
- 2. Brick--glazed and unglazed
- 3. Transite
- 4. Wood
- 5. Metal
- 6. Conducting non-sparking floor

The approach that was taken to the development of these sampling protocols involved the following steps:

- A. Literature Search
- B. Sampling Protocol Selection
- C. Analytical Method Selection/Development
- D. Certification Testing
- E. Development of a Spike and Recovery Test Plan
- F. Spike and Recovery Testing
- G. Interference Testing

The <u>Literature Search</u> provided an overview of existing sampling and analysis methods. The results of the literature search suggested that most existing methods would not satisfy the Army's requirements as stated above. Thus, in the Sampling <u>Protocol Selection</u> step, several approaches based on technology developed for applications other than explosives analysis were proposed for further development. Those approaches involved methods intended specifically for qualitative analysis, and a method intended specifically for quantitative analysis of explosives/explosive residues on surfaces.

Development of the quantitative analysis method proceeded with an Analytical Method Selection/Davelopment step, in which advantage was taken of the fact that several methods for the dtermination of explosives were present in the USATHANA data base. These methods were modified as necessary to permit their application to the present study, and the precision and accuracy for each of the resulting methods were assessed in preliminary Certification Testing (semiquantitative).

Subsequent to approval by the Technical Project Officer of a Spike and Recovery Test Plan, methods for each analyte on each surface type were evaluated by Spike and Recovery Testing. Finally, methods were subjected to Quantitative Certification Testing according to the requirements in the 1980 USATHAMA QA Plan. Since most of the analytes are determined in a single extract, the need for Interference Testing in the laboratory was largely eliminated, interferences from other sources which may be present in samples collected in the field were not addressed in this study.

This study resulted in the development of several methods for the sampling and analysis of explosives/explosive residues on building materials surfaces. A method for qualitative determination based on detection of charge-transfer complexes formed between the explosive/explosive residue and a visualization reagent applied to the surface has been evaluated in the field at two army ammunition plants. A method for quantitative determination based on solvent extraction of samples collected in the field followed by high pressure liquid chromatographic analysis has been evaluated on samples prepared in the laboratory, and found to give promising results. The theoretical and practical feasibility of another method for qualitative determination based on UV irradiation of a suspected contaminated surface with subsequent detection of any explosives/explosive residues present has been demonstrated. This approach may provide the means for "scanning" an area to determine whether explosives are present on a real-time basis. Further development of this approach is recommended. No methods for inorganic species which would represent substantive improvement over existing qualitative and quantitative methods were identified.

Detailed discussions of these areas of investigation are presented in the remaining sections of this report.

II. SAMPLING AND ANALYSIS METHODS SELECTION

A. INTRODUCTION:

The available literature on the sampling and chemical analysis of explosives/explosive residues was reviewed to provide an overview of existing sampling and analysis methods prior to selection of specific approaches for further development. The findings of that review are described below.

B. LITERATURE SEARCH:

Work was initiated by conducting a literature search to identify existing sampling and analytical methods for explosives which might be applicable to the particular problem of detecting and determining those same materials on building materials surfaces.

1. Approach.

There were basically two problems that had to be resolved in order to develop appropriate strategies for the computerized literature searchings: (1) if the search strategy involved the fifteen defined analytes in combination with the defined surfaces, as well as with general terms such as surface sampling, recovery, sample selection, detection, etc., the search results were nil; and (2) if the search strategy involved the use of such general terms as explosives, ammunition, and dynamite, as well as the Chemical Abstracts Section 50 (Explosives and Propellants), combined with terms such as analysis, sampling, recovery, identification, determination, etc., the search results were not applicable to this study.

The techniques used in the forensic sciences appeared appropriate for the recovery and analysis of the analytes under consideration; therefore, the search strategy that yielded the most relevant citations involved the combination of the <u>Chemical Abstracts</u> Forensic Analysis Subsection, the Registry Numbers of the 15 analytes, and the terms explosives, dynamite, and ammunition.

2. Sources and Coverage.

The search for applicable techniques for surface sampling for explosives/explosive by-products has been performed on:

Source Coverage Number of Citations

Chemical Abstracts 1967 - present 164

The 164 citations were reviewed for duplication and extraneous material and were reduced to an output of 45 citations. This output was reviewed and the 45 citations for document retrieval.

Arthur D Little Inc

In addition to the computerized search of Chemical Abstracts, a manual review of a National Technical Information Service (NTIS) bibliography with abstracts, entitled Pollution Caused by Ammunition Manufacturing, was undertaken. Of 237 abstracts, 9 appeared appropriate for review and were ordered.

The following computerized data bases were also searched:

APTIC
ENVIROLINE
POLLUTION ABSTRACTS
SCISEARCH
NTIS
CRIS (Current Research Information Systems)
SSIE Current Research
CONFERENCE PAPERS INDEX

Results.

The results of the literature search are summarized in Tables II-1 through II-4. Table II-1 is a bibliography of useful citations including (1) the authors' abstracts or brief summaries taken directly from the article, describing the purpose and results of the work described in the article, and (2) comments describing potential applications of the work described in each article to this study; Table II-2 lists the analytes of interest in this study for which sampling and/or analysis procedures are described in each article with detection limits, when included in the article, listed under the respective headings in the table; Table II-3 lists the analytical methods used or recommended in each article for determination of analytes of interest in this study; Table II-4 lists (1) the sampling procedures used or recommended in each article for analytes of interest in this study, (2) any additional sample preparation procedures required subsequent to sampling and prior to analysis, and (3) any interferences described in the article with the sampling and/or analytical procedures.

TABLE II-1. BIBLIOCRAPHY OF USEFUL CITATIONS

. . Specific colorimetric test for RDX; would require meparation of sample from surface; 100°C best source required; which may be difficult

in field applications.

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percendings. This paper describes a new approach to the their libraries of mach eshibits. The sector control control is described the presence or absence of me man to penaltie of the comments and legections. be about chestral tests done so a spec plate. The croality see the compared with the manufacturer's data and is sent cases, it is found that there is said one employees with the combination of improvious: Navelibed, a full Measificaton of a sample automass other compounds, in particular wegats and aldehydes; The detection and identification of altro-bodies is a common requirement in the explosives inducatory. The matitize of a 5 to 10-mg specimen of one drop of accremental man and drop of retrancial comments on the residual provides a bloc reloc with district observe and a dark red with trinitratologue. The test has been applied to a wide range of obsider and current industrial blassing explosives and it has been found that man of the other compresses presents present interfere of the fearther. Supplies of explosite frank of a series have often The familian spot test for the identification of cycletimethylemetrisicramine, RPA, is based on compound in the presente of thomas and altropen-free sulpharic acid. As objection to this test is, becaver, that a rel colour is preduced by determinated and tend to be small, margineeur tive and concentrated with material from the the production of the rad colour formed by this malpharic acid is of the quality marketed as "migrogen feet." A method of auranumiting this a deer plat colour may be produced aniess the difficulty has been forme. AUTHOR ABSTRACT The ideactfiles the of immerital Interrital Standing Explosives. rintermier. Analyse (9):5/8; Amos, S.A.M.; Yallop, H.J. A Test for Cycletrimethylene-Masters: 91(51:336-137; 1966. Blacking Systemiers of the Gellerite Tope, J. Farmalic Amas, S.A.H.; Tallop, B.J. The Detection of Dinitro and Triaitre Areastic Redies in F. 121/41:15 . 15 . 15 Acess, S.A.M.; Talles, B.J. REPUBLICE ARTICLE MUMPHER

test for 104 deep max toquire leating; later-May be maeful as mpot test for 2,4-DMT and 2,4,6-TMT directly on surface; interferences foregree free explosives of types other than these listed and discussed. May be marful as spot test for 2,5-DHT, 2,6-BHT, 2,5,6-THT, MT, MTE, MINT wand for detection of 2,6- and 2,6-BHT and 304 are Mercy factorized to those to Merc. 1 and 2, except from explosive types other than aftrocompanies on discussed.

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TABLE 11-1. BIBLIOCANNET OF REEPIL CITATIONS (CONTINUED)

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Canifold, B.V.; Deforement, P.E.
The Ther of the Condolff Camera as a Servening and Confirmation Tool
in the Amalouis of Explosive
Ecolomes Chambler, C.D.; Kobbbech, J.A.; Bolleter, W.T. Costismoss DT Freess Studies, III. This-Layer Gressrographic Analysis of Oxidation Products from Mitterfom J. Gremstogr. 64:123-126; 1972. Chacan, D.E.; Morvitz, G. Qualitative Analysis of Prisers, Tracers, igniters, incendiaries, Bacters, and Delay Compositions on a Microscale by Use of infrared Spectroscopy, Microchem. J. 17: 31-60:1972

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This study pressures a scheme for systematic analysis of emploater residunt isolated by physical tensoral or soluter extraction. The techniques are a combination of infrared spectroscopy, TL, I-car differentian, unisolan spectroscopy, alcreaces.

As ideal technique would have to be bath secultive and definitive and, is addition, should be able to deal dethints are contained to be subjected with actual explorative residue amples. This ideal with actual explorative residue amples. This ideal with actually approximately in create, which is capable of producing a detailed I-ray diffraction pattern for a single microscopic crystal. Contamination difficulties can be chalminated if the requisite intact microscopic crystals can be found in the residue.

Excepting, merhanically removing, and identifying these crystals climinates the meed for a preliminary polynet extraction with its attentiant contains and manufacturing problem.

This-layer chromatograph meparations are shown of all major oxidation products from various aterof the continuous THT mitration process and of mitration compounds resulting from inpurities in the tolucus used for mitration. This laboratory undertook an investigation on the application of infrarel spectroscopy to the detection of the organic compounds and invegation compounds preprint in primers, tracers, igniters, incendiaries, boosters, and delay cumpositions.

Or combitions given for determination of MFT and tribits demand from to 9.4001 in IMT; fill detector used; mealings performed on became entract of TMT nample.

THE and NTB acced on prime identification techniques for high explosive with other materials mand for confirmation. Analyses performed on other and arrives errivers from other, used, and ecoling tile. Columnation team inclination of the firm term in the second of the columnation of the columnation of the second of the columnation of the columnation

Probably set useful is this study due to smannel testrosestation requirement. TLC conditions for nitro-compounds.
Visualization with UV light and subsequent
colorimetric development.

frohably not uneful in this study except as screening or confirmatory technique. Relatively large sample size (1.e., 1-2 mg) requirements.

COMMENTS	Collection of analytic vapors may provide a means of eliminating interferences encountered in solvent extraction sampling methods. Quantitative capabilities of method for determination of analytes at extended time after deposition uncertain.	Non-specific test for nitrate. Prohably not useful in this study due to large numbers of potential interferences.	Conditions given for separation of synthetic mixtures containing various analytes of interest on Corasii il column, UV detector.	The conditions for separation of various explosives. Visualization method not specified.	The conditions for separation of various explosives. Uses consercially available portable TLC kit and battery-povered by lamp for vizual-station of resolved constituents. May be adapted for this study if conditions for resolution and visualization of other analytes of interest can be demonstrated.
AUTIOR ABSTRACT	This paper describes the use of a short column containing porous polymer beads to collect explosive vapors of ethylene glycol dinitrate (RibN), nitro-glycerine (NC), and trinitrotolucue (TNF), vitn subsequent analysis by TCL.	It is (the author's) opinion that a frank discussion and demonstration of the fallability of the so-called "Paraffin Test" would be appropriate Attention has been directed to evaluating reagents, improving techniques of casting and application of reagent, so well as devising a satisfactory manner of revoiding the reactions.	The mualysis of energetic compounds is often complicated by their low volstility and low thermal stability. A potentially useful technique for such analyses is high speed liquid chromatography since it does not require sample heating or vaporization. The use of the method is illustrated using synthetic mixtures of components found in explosive formulations. Separations involving nitrate esters, infrocramatic compounds and nitramines are discussed. The use of the technique for the analysis of single and double base propellants is also presenting.	In complicated cases of explosive mixtures, degraded explosives, explosives containing copolymerised planticisers, or with impurities present, the circumacographic separation detailed in (a previous article by these authors) may be inconclusive. Furthermore, specific chromatographic separation may be required to detect the explosive constituents. [In this article] the authors describe how to seat for various nitro esters.	In this work the approach was to develop a field kit that incorporated a nondestructive uniform analytical technique with emphasia on protability, east of operation, and low cost. The report details the modification of a commercially available portable kit for the achievement of these objectives.
REFRENCE	Chrostinaski, J.R.; Holmes, R.N.; Rehn, B.W. The Collection and Determination of Ethylene Clycol Dinfrate, Nitroglycerline, and Trinitiotoluene Explosive Vapors. J. Forensic Sci. 21(3):611-615; 1976.	Cowan, M.E.; Purdon, P.L. A Study of the "Paraffin Test." J. Forensie Sci. 12(1):19-36; 1967.	Donli, J.O.; Juhasz, A.A. Application of High Speed Liquid Chromatorraphy to the Qualitative Analysis of Compounds of Propellant and Explosives Interest. J. Chromatogr. Sci. 12(1):51-56; 1974.	Ellu-Colmet, J.; Forestier, H. Characterization of Explosives' Traces After an Explosion. Int. Crim. Police Rev. Series 325:38- 47; 1979.	Fisco, N. A Portable Explosives Identification Kit for Field Dsc. J. Forensic Sci. 20:141-147; 1975.
ARTICLE	<u>e</u>	z.	13	=	2

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SLIKHARO	Useful for metals only.	Useful for clemental analymie only. not useful in this study due to unus instrumentation requirement.	Sample collection methods used for d of metals (Es and Sb), and may not b (except for cotton swab technique) f nation of organic compounds.	Summary article including recommendation of the sumpling and analysis procedescribed is other references. Biolicechnique not discussed in detail, howerth further investigation.
AUTHOR ABSTRACT	Pb and Sb, present in common commercial firearm detocators, were determined by anothe stripping polarography on the hands of persons having fired revolvers. The hands were cleaned which filter paper mojetesed with 1800, and the ashed paper was analyzed in a commercial apparatus.	The deposits resulting from the discharge of 0.22 calibre summittion have been studied by neutron activation analysis and suterading raphy.	Of the various collection systems currently in use, only cotton means and film-lifting proceduren appeared to combine came of use in the field with suitability for rapid laboratory examination by FAAS. However, one additional material, transperent adhesive tape, also appeared suitable for routine use. This material is readily available, inexpensive, convenient to use, and amenable or reproducible nample collection.	Stoluminescence may be uneful for detecting employers traces. In this technique, special cultures are negatived to vapors icm a specific explosive and respond to the presence of this explosive by the emission of light. While there
REFERENCE	Gagilano Candela, R.; Colemas, M.; Strada, L. Identification of the Explosion Residues of Shots from Firearms by Anodic Stripping Polar- ography. Zacchia 12(1):44-52; 1976. CAS Mr. 85:1546485.	Gislasos, J.; Pate, B.D. Studies of Camshot Residue. J. Radio- analytical Chem. 15:103-113; 1973.	Goleb, J.A.; Hidhiff, C.R. Firearms Machange Kesidue Samule Collection Techniques. J. Forentic Sci. 20: 20:-707; 1975.	Hoffman, C.M.; Byall, E.E. Identi- fication of Explorive Residues in Ecub Scene Investigations. J. Forestic Sci. 19:54-63; 1974.
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determination be applicable for determi-

Probably

Jumple Scene detail, but my be lations of countries

technique not discussed in di worth further investigation. mescence units are compact, portable, relatively imagemative, and do not require extensive operator Particles of the explosive eved in a bonk can the lengthy activation time of the culture when a new detector cell and be prepared, the bioluni-

removing traces of vortions explorings from the bonds of a subject who has recently handled them. Exploring Cotton made maded in acatemy are effective in Total a in mat ca a tentille to it seburpase trate of the estreet. Idea xore by T.C Preceders.

sessittatry seconsary to actect the traces of explusive debris may contain substances that would interfere with present in a large malme of debris. In addition, the

Ideatified by rather simple chemical and imitramental

procedures. As a last resort, solwest cuttactive of

the debrits may be performed, but this is not recon-

araded stare it usually lacks the specificity and

takes at the scene of the debuts is meticulously examined micraecopically. The explosive can then be

usually be recovered from properly collected debris

Introduces such as the fill have tolar in caldly autholice for the presence of explusions. The explosive Scanning debris, cectes hand made, clicking, and air present in the confirmative changes we thank.

NT I CLE	Denois	ABTIOR AKSTRACT	SICOMAGO
P	Hoffnemer, J. Quantituitys Analysis of Mitro Conpounds in the Micro- in Ficegram Empe by a Combinetton of Thire-Layer and Vapor Phase Chrum-tography With the Michel-6, Electron Capture Metertor, J. Chromotog, 51; 243-23; 1570.	A method has been developed for the quantitative analysis of mittee compounds in the micro- to picugius range by a combination of this-layer and vaper phase chromatography employing the mittel-by electron capture detector. The relative electron absorptivities of ten mitte compound, have been measured with 1,35-trimitrobenius as a standard. Experimental variables affecting the mithel-by election capture detector are presented.	Conditions to separation of altro- compounds by TLL, visualization by UV with set ilents :0.6 pg/spet; spots reserved fr a TLC plate by suppling and then entracted; altro compounds in ex- tracts quantitated by CC with RI-6, detector with recoveries :952 for com- pounds tested; disadvantage is that RI-6, detector is easily consadanted or overleaded.
R.	Jenkins, R.; Fallos, H.J. The Identification of Explosives in Trace Quantities on Objects Bear an Explosion. Explosivatoffs Re. 6:139-141; 1970.	A [TLC] tecimique is described whereby traces of explosive affecting to sucremaling objects after an explosion can be detected and identified. The method has also here applied to the identification of traces on the skill and clothing of persons numbered to have been in contact with explosives.	The conditions given for aeparation of various explosives: colorisation visualization,
ă	Jones, P.7.; Membler, R.S. A Netolumi erscence Technique for Deroction of Gammary Residue. J. Forenaic Sci. 20(2):231-242; 1975.	In our study we concentrated on the development of a unlecular photolominencence technique to detect metallic elements in granulor residue. As discussed subsequently, the use of this technique is attractive because of the case of analysis and its semilifyity and low cost. In this prilinlancy study we were concerned with the detection of Pb. Sb. and En.	Resful for metals unly.
;;	Kaples, M.A.; Zittis, S. Identi- fication of Post-Explosion Residens. J.A.D.A.C. 60(3):623-624.	A scheme for the identification of explosive residues from post-explosion scenes is described. The first step consists of organic and inorganic extractions rather than microscopic examination. The methods of identification which follow include thin layer and gas-liquid chromatography, infrared and ultraviolet spectroscopy, and chemical tests. The system, which has been used routinely by the larael Police for 5 years, deals efficiently with atambard military explosives as well as with home-made, improvised mixtures.	Conditions given for various methods used for qualitative identification.
53	Kelsvil, J.F.; Burton, R. Vari- ability in the Chemical Contum- institute Effects of Cam Shot. J. Radicanal, Chem. 31:45459; 1976.	This is a contribution to a larger study alacel at develop- ment of a technique to determine the origins of waterfoul from their feather chemistry, using automated X-ray fluc- trescence spectrometry. Since feather samples commonly come from whot birds, an effort was made to measure the contamination effects of whot using corton cloth to simulate feathers. At pointblenk ranges contamination can include the elements Bs, Sb, Pb, Cu, S and likely others depending on the exact componition of both gun powder and shot.	Useful for metala only.
2.4	Kempe, C.R.; Tannert, W.K. Detection of Dynamic Residues on the Bands of Bombing Suspects. J. Forensic Sci. 12(2):323-324; 1972.	Studies were undertaken mains TLC for the identification of nitrate estern such as nitroglycerion, claylone elycoldinitrate, and pentacrythritor retranitrate (PRTM). An strempt was made to show the reliability of TLC at means of identification of these nitrated estern when emayed from the hands.	TIC couditions given for separation of various explosives; colorimetric visualization.

COMMENTS	Useful for metals only. Sample obtained by vashing hand in IM HCI.	Useful for metale only. Sumple obtained by washing hand in 1M HMO3.		Useful for metals only. Sample obtained by washing hand in 1) detergent, 2) 0.5 H HCl.	Useful f metalo only.
AUTHOR ABSTRACT	In this study conditions for the determination of lead and entimony in gunshut residues by anodic stripping voltammetry using a mercury-coated graphite electrode are established. A cample is collected by washing the hand in 1H hydrochloric acid. Lead is determined in a portion of this sample, with 1H hydrochloric acid as electrolyte. Antimony is determined in a second portion, with 4H hydrochloric acid as electrolyte. Prior separation or preconcentration steps are not required for either analysis. The procedure has been applied to samples obtained from five "normal" hands after they have fixed a weapon.	Because the amounts of the elements analyzed are at microgram levelo, specially designed procedures and training are required to be able to collect the samples without contamination. The existing techniques, such as paraffin lift and colton swabbing, are found to be unsatisfactory in this respect. By taking repeated samples from hands by these procedures, it was found that three or four collections are required for complete removal of the trace elements. Thus, a single collection by these methods is not quantitative, and therefore, any subsequent calculation would be in error.	Neutron activation analysis is not effective in detecting lead, which is one of the important constituents of leakage residues. Hence, a method such as atomic absorption spectrometry (AAS) must be used in addition to MAA for this analysis.	Non-destructive analysis of gunshot residuce on bands was studied by anotic stripping voltametry using a low-cost home-constructed polarograph. The washing solution was 0.5M HCl. Antimony was first snalyzed in 4M HCl solution; and zinc, lead and copper were then similtaneously analyzed in 0.2 M sectate buffer (pH 5.9 ±0.1).	An anodic stripping procedure was developed for the simultaneous determination of a zinc, cadmium, lead, antimony and copper mixture, and used for testing the instrument. Effects of pre-electrolysis time, concentration changes, presence of other species and overlapping were also studied.
REFERENCE	Konanuc, M.K.; vanloom, G.W. Determination of Lead and Antimony in Firearm Discherge Residues by Anolic Stripping Voltammetry. Talenta 24:184-187; 1977.	Krishnan, S.S. Detection of Gunshot Residue on the lands by Neutron Activation and Atomic Absorption Analysis. J. Forensic Sci. 19(4): 789-797; 1974.		Liu, J.N.; Lin, W-F.; Nical, J.D. The Application of Anodic Stripping Voltametry to Foreusic Science. II. Anodic Stripping Voltammetric Analysis of Gunshot Residues. Forensic Sci. Intl. 16:53-62; 1980.	Liu, J.H.; Taylor, L. The Application of Anodic Stripping Voltammetry to Forensic Science. Forensic Sci. Intl. 16:43-52; 1980.
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REFERENCE

Libyd, J.B. Detection of Microgram Amenits of Nitroglycerin and Related Compounds. J. Forenste Sci. Soc. 7:

Systematic Approach to the Detection of Explosive Residues. III. Commercial Dynamite. J. A.O.A.C. 57 (5):1092-1097; 1974. Midkiff, C.R.; Washington, W.D.

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Systematic Approach to the Detection of Explosive Residues. IV. Military Explosives. J. A.O.A.C. 53(6):1357-Hidkiff, C.R.; Washington, W.D. 1974; 1976.

Parthar, D.B.; Sharma, S.P.; Verma, K.K. Trace Analysis of Ex-plosived as Pi Complexes. J. Forenste Scl. 13(2):246-252; 1968.

AUTHOR ABSTRACT

Nitrate enters, on hydrolysis, pactly disproportionreaction. The test may be adapted to the characterof nitroglycerin in air and biological materials, the nitrous acid formed being detected by Griesa'u ate to carbonyl compounds and nitrous acid. This has provided the basis of tests for trace amounts Ization of explosive components in very small samples such as fingernall scropings.

and ethylene glycol dinitrate, which are present in most connected dynamites. Problems encountered with Additional chemical tests performed on the debris to facilitate identification of the particular type of discussion of major categories and typical formulagraphic and infrared techniques are utilized for the identification of the explosive oils nitroglycerin tions of dynamite is included. Thin layer chromato-Tests are described for the detection and identification of suspected dynamite in bombing debris. A solvent extraction of these oils are discussed. dymalte present in the debris are described.

2,4,6-trinitrotoluene, cyclotrimethylenetrinitrumine, for the ideinfication of major explosive components. Tests for minor components to enable characterization collected at the acene of a criminal bombing. Major and to distinguish similar compositions are included. detection and identification of military explosiven categories and typical formulations of some common military explosives are described. Tests are deschromatography and infrared spectroscopy are used Part IV of the series encompasses tests for the cribed and evaluated for the identification of and pentaerythritol tetranitrate. Thin layer

amlans. The present paper describes a simple and con- visualization. vesteur method for the identification of trace amounts of explosives as charge-transfer complexes with amines chromatoplates. It was possible to identify as little In the detection of explosive advantage may be taken of their highly colored a complexes with acomatic being highly culured could be located enaily on the 18 1-2 micrograms of an explosive as its a complex. employing T.C technique. The resulved a complexes

TLC conditions for separation and detection of nitrogiyeerin in presence of nitrate esters. Colorimetric visualization.

Colorimetric Visualization, Qualitative iden-tification confirmed by other methods (optical TLC conditions for separation and identification of nitroglycerin in dynamite residues. microscopy, GC, 1R).

residues. Colorimetric visualization. Qualimethods (optical microscopy, CC, 1R, XRD); spot tests for INT and RDX performed directly on polyeuter adhesive tape used to collect TLC conditions for separation and identification of TNT, DNT, RDX and PETN in explosive tative identification confirmed by other particles.

The conditions for separation and identification of various explosives. Colorinetric

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CVERNITS	has recently TLE used for confirmation of MBAN in presence manufacturing of various ofner explosives. Colorimetric replacing visualization. Gredients are ne nitrate outcoured with for the Inthis paper. In this paper	ists on a select Sput tests for several explosives of interest. Hund in explu- licating rut required for most tests. Tests ited. Any un- may be performed on visible particles or hysical exam- solvent extracts. Confirmation by TLC, IR, yen here.	sts used by TLC conditions for confirmation of qualitative blosives and identification of several explosives of int 2 presents terest. Colorimetric visualization. Ethods used in commonly used	onfirm the TLC conditions for identification of NC in sucked as powder flakes. Physical transfer of sucked is visible particles, transfer tape, and culture olubility in extroction used for sampling. They are supplice plates, transfer to the sucked for sampling.	ims covering Probably not useful in this study except as I many possible confirmatory technique due to 1) relatively elated com- large sample size requirement, 2) can be applied only to pure compounds.	thed finitic. Spot test for lead. Cotton swabs moistened Therefore, with 25% acetic acid used to obtain samples ie help in from gun barrels.	n foreusic Useful for metals only. I integrated with gension liny, emission
AUTHOR ABSTRACT	The F.I. dufont de Nemours 6 Co., Inc., has recently announced that it is discontinuing the manufacturing of nitroglycerin-based dynamites and is replacing them with formulations whose primary ingredients are assentium nitrate (AN) and monomethylamine nitrate (PPAN). The foreusix chemist is thus confronted with the need to be able to analyze realdues for the possible use of this type of explosive. In this paper procedures are given for the qualitative determination of MMAN.	In this paper the results of various tests on a select number of ione and organic compounds found in explu- sives and explusive regidues are presented. Any un- explosed explusive particles found by physical expa- ination can be tested by the methods given here.	In Part 1 of this paper the chemical tests used by this laboratory in the screening of explosives and explosive residues were discussed. Part 2 presents the thin-layer chromatographic (TLC) methods used in this laboratory for the confirmation of commonly used organic explosive compounds.	A simple and inexpensive procedure to confirm the identity of unburned or partially burned flakes of smokeless powder is described. The procedure is based on (1) particle morphology and solubility in acetone, (2) R _f values of the flakes when they are ciromatographed on thin-layer chromatographic plates, and (3) specificity of the visualizing reagent to nitrite.	A compliation of 68 infrared spectrograms covering all common high-explosive compounds and many possible explosive ingredients, additives, and related compounds has been prepared.	Metallic traces are expected to be retained instact the barrel even after usual cleaning. Therefore, the metallic traces can provide valuable help in determining whether the firearm had ever been fired.	The examination of gunshot residues in a forensic science laboratory should be a series of integrated procedures. In this paper we will deal with gunshot residues utilizing soft X-ray radiography, emission spectroscopy (ES), and AS.
REFERENCE	Parker, R.C. Analysis of Explosives and Explosive Residues, Part 3: Honomethylamine Nitrate. J. Forensic Sci. 20(2):257-260;1975.	Farker, R.C.; Stephenson, M.O.; McOwen, J.H.: Cherolis, J.A. Antaysis of Explosives and Explosive Revidues. Part 1: Chemical Tests. J. Forendic Sci. 20(1):133-140; 1975.	Furker, R.G.; McOven, J.M.; Cherolls, J.A. Anslysis of Explo- sive Residues. Fart 2: Thin-Layer Chromitography. J. Forensic Sci. 2012):254-256; 1975.	Peak, S.A. A Diln-Layer Chromatographic Procedure for Confirming the tresence and Identity of Smokelies Powder Flakes. J. Forensic Sci. 25(3):679-681; 1980.	Pristera, F.; Halik, M.; Castelli, A.; Fredericks, W. Analysts of Faplosives Using Infrared Spertruscopy. Anal. Chem. 12(4):495-508; 1960.	Stuba, J.K.; Migra, G.J. Projectile Traces. Int. Crim. Police Rev. Series 288:150-151; 1975.	Stone, 1.C.; Fetty, C.S. Examination of Gaushot Residues. J. Foreusic Set. 19(4):784-788; 1914.
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S.D.G.W.D.C.	HPLC conditions for separation of NG, RDX, TNT from mixtures. Confirmation by chemical loni- sation mans spectrometry.	Recommended approach to identification of trace quantities of homemade, commercial, and mill-tary explosives and incendiaries involves 1) optical examination of debris under low magnification for detection of unburned explosive residues, followed by 2) solvent extraction.	Detection of vapors by Vapor Trace Analyzer (VTA) manufactured by Hydronautics-largel Ltd.; probably not useful in this study except as ecreening technique because of likely difficulties in quantitation.	Recommendation for swabbing of surfaces.	TLC conditions for several explosives ordinarily occurring as impurities in TNT; colorimetric visualization.	C1-FE MS of pure compounds and residues; probably not useful in this study except as sereculng or confirmatory technique.
AITHE ADSTRACT	High performance liquid chromatography and chemical mans spectrometry have been applied to the isolation and identification of explosives. The use of amounta as a reagent gas for chemical lonization has been evaluated and its advantages ever methane, water, hydrogen, and isobutane are discussed on the basis of data from common explosives. The off-line LC-MS approach has been applied to the analysis of simple residues from test explosions under controlled conditions to simulate an actual bombing.	An analysis scheme, in several parts, dealing with the identification of explosives and incendiary deatructive devices encountered in actual cases is described. Part I is introductory and is the first step taken in the examination of bomb debris for the various types of explosives and/or incendiaries. Examples and photographs of typical bomb residues are presented.	A Vapor Trace Analyzer (VTA) was found to be a valuable tool for scanning bomb debris for traces of certain types of explosives. The application of the VTA in screening bomb debris and locating secreted explosives and uses in other types of physical evidence is discussed. Thin layer chromatography is used to identify physically removed unspected explosive particles and samples from solven extractions of blast material.	The use of explosives in breaking offences, and in some techniques in the investigation of their use is described.	This article describes a two-dimensional TLC method for the separation and identification of u-TNT impurities, including some oxidation-reduction products of decomposition as well as common production grade impurities. In addition, a unique detection method is described in which the reductor of the developing reagent is directly incorporated in the thin layer.	The value of CI-MS in combination with EI-MS has been demonstrated as an analytical method for the identification of forensic compounds. Duta acquisition consists of converting the recorded mass spectra into plotted and tabulated normalized mass spectra into plotted and tabulated normalized mass spectra by using a central computer. Chemical ionization mass spectral library comparison and identification are done manually.
30M a 139 a	Youro, P., Peteraen, B.A., Colwell, L.; Karger, B.L.; Harria, U. Anniysia of Kaplosives by High Performance Liquid Chromatography and Chewical lonization Mass Spectrometry. Anal. Chem. 49(7): 1039-1044; 1977.	Mishington, W.D.; Midkiff, C.R. Systematic Approach to the Detection of Explosive Residues. 1. Basic Techniques. J. A.O.A.C. 55(4): 311-822; 1975.	Washington, W.D.; Midklff, C.R. Systematic Approach to the Detection of Explosive Residues. J. A.O.A.C. 56(5):1239-	Vallop, II.J. Breaking Offences with Explosives—The Techniques of the Criminal and the Scientist. J. Forensic Sci. 14:99-102; 1974.	Vasuda, S.K. Identification of laportities in a-Trinitrotoluene by Thin-Layer Chrowatography. J. Chromatogr. 13:78-82; 1964.	Vimon, J.; Zitrin, S. Processing and interpreting Mass Spectral hata in Forensic Identification of Drugs and Explosives. J. Forensic Sci. 22(4):742-747; 1977.
ARTICLE	0,	7	3	÷.	55	4

COPPENTS	Useful for metals only.	Spot teets for TNT and RDX. Advantage is that tests were performed and evaluated directly on surfaces of interest.
AUTHOR ABSTRACT	The use of the Weisz ring oven to localize, concentrate, separate, and identify the four actals of interest from each other and from possible interfering fond and other extraneous matter, in described.	As explosive detection spray system has been deviloped which will detect explosive residues on the exterior of package and letter bombs. The detection is through the formation of colored neutron products using selected spray reagents.
REFERENCE	Bosen, S.F.; Scheuing, D.R. A Rupid Microtechnique for the Detec- tion of Trace Metals from Gunshot Residues. J. Forensic Sci. 21(1): 161-170, 1976.	Mywnt, R.E. Development of a Simple Fortable Detection Kit for Selected Explosives. D.T.L.S. Report No. TR-185, September 1977.
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TABLE 11-2. ANALYTES DETERMINED IN LISTED CITATIONS

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TABLE II-3. ANALYTICAL METHODS DESCRIBED IN LISTED CITATIONS

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TABLE IL-4. SAMPLING AND SAMPLE PREPARATION PROCEDURES DESCRIBED IN LISTED CITATIONS

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	Article No.	11 2	TOTAL TRACEDORES DESCRIBED IN LISTED CITATIONS	LISTED CITATIONS
	-	Standard samples.	Sample Preparation	Interference
			Mix sumply one of contract	
			leaf	Pale bule-green color of cyclo- tetramethylenetetranitramine (HMX) may be cathigufehed from
	7	Standard sa a vies.		Aux by repeating test at 150°C; RDX is the only ampd. commonly found in explosive formulations that Rives a blue celor in this test.
			Mix sample and rgts.	In the prosence of MG, color produced with DMT is prosen
	•	Cutton wool swabs soaked with ether, water,	;	Initial color should be observed since changes occur with time.
	7	Sellotape transpacent adhesive tape used for physical transfer of particles from skin,	Mix sample and rgts. Nefer	
	ĸ	Extraction with hexane,		
1	ع	Extract residue with ether, acetone.	·	
18	,	Physical transfer of residue crystal from auritan	Spot sample Aoln, on TLC plate, develop,	
	æ	Actions solution of process sample.	None.	
	5	Standard samples.		
	2	Callect vapors on short column containing porous polymer beads.	KBr pellet. ods. Extract column with account	
	Ξ	Paraffly cast of bands of persona known or suspected to have discharged flrearms.		
	12	Synthetic mixtures.	Mix rits.	M) _j -1on
	1.3	Not sportfled,		
Art	21	Physical transfer of explosive fragments from surface.	Spot sample solu, on TLC plate,	
hur	g g	Hands of persons known to have discharged fittearms were eleganed	election ample solu, en TLC plate, develop.	
ז ח	3	Surface sample analyzed diverrly,	Ash (Uter; dissalve residue,	
irela		Coffeen south, film—lifting with films made from solms, of film-forming polymers, transparent adhesive tape.		
Inc	JB P	Enysteal transfer of explosive fragments from sojface, solvent extraction, coffee-solvent		

TABLE II-4. SAMPLING AND SAMPLE PREPARATION PROCEDURES DESCR

		STATE INFORMATION PROCEDURES DESCRIBED IN LISTED CITATIONS (CONTINUED)	_
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19	Standard samples.	Sample Preparation Interferences	<u>.</u>
		ite, id ed,	
54	Solvent extraction; cotton v. I swabs soakded in appropriate solvent used for extraction of compounds from bonds on a con-	resulting soin, ahalyzed by GC.	
ភ		Spot sample on TLC plate, develop.	
22			
2.1	Direct manysis,	Method dependent .	
24	Cotton swabs soaked in acetone.		
25	Wash isand to 1 M nos	Spot sample soln, on TLC plate, develop.	
56	Wash trand to 1 M HMM.	Codmilum	
7.2			
į			
28	Standard samples.		
82	Plassolve motoring to the property of the prop	Aid O.1 M acreate buffes to sumple soln.	
	and the sections.	Spot sample Boln. on TLC plate,	
홌	Extraction with chloroform, acetome.	develop.	
=		Spot sample solu, on TLC plate, develop.	
	motstened with chlorof acetone; for hands, cotton wahs	Spot sample soln, on TLC plate,	
71.	Standard samples in acetone solution.	· document	
Ξ		Spot sample solu, on TLC plate, develop.	
	scindid soles. In witer.	Spot sample solu, on TR plate,	
%	Extraction with acetone.	devertable.	
		Mix sample and tyts, Extractions material.	-
		Substitute methane or ether for account in extraction step or take up residue from accione	or take
ú	Extract but with professor	extraction of the these solvents.	311
		Spot sample solo, on TIC plate, develon.	

TABLE 11-4. SAMPLING AND SAMPLE PREPARATION PROCEDURES DESCRIBED IN LISTED CITATIONS (CONTINUED)

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Sample Preparetion	Place porfile or spot ample soln.		A						Spet maple wie, as til plate,		Place nample mets, on filter expert, week, mass case and and	
Samiling Procedure	Physical transfer of visible porfiles, transfer tape, or culture entraction.	Standard sumples.	Catter souts motatemed with 35% acette acid.	for eloth, extractive with Many; for heads, wanding with Many, Mcl.	Simplified samples; oceans extraction.	Optical examination under les mapeification, polyent extraction.	Defection of vapors.	Seath.	Standard samples in chiesewhern.	Mire: mairely of residue detained after competition of solvent from actions extinct.	It is activation	Spene respents directly as anythmy.
Article Ib.	£	17	ĸ	£.	5	7	;	5	3	¥	1	12 2

C. REVIEW OF SPOT TEST CHARACTERISTICS:

At the request of the Technical Project Officer, a review of the spot tests used for detection of explosives/explosive residues was carried out concurrent with the literature search to determine whether those tests could be used for in-situ detection of explosives on the surface types of interest in this study.

Tables II-5 through II-8 list and describe selected characteristics of spot tests which may affect their suitability for use for qualitative analysis of explosives/explosive residues on the surface types of interest. Tables II-5 and II-6 list spot test methods and characteristics by analyte. Tables II-7 and II-8 list spot test methods by reagents used in the respective methods, and list and describe selected properties and characteristics which may be exhibited by those reagents when used under the conditions specified in the method.

To evaluate the data included in these tables, we defined a set of minimum criteria which a spot test proposed for use for the contemplated qualitative analysis should satisfy. These criteria were: (1) the spot test should permit rapid screening of large areas; (2) it should be sensitive down to the agreed upon detection limit of 0.5 µg of analyte/cm² under various surface conditions; (3) it should be specific for the analyte of interest; (4) it should not result in an irreversible chemical reaction which would make the analyte unavailable for subsequent quantitative testing; (5) its use should not present an unusual hazard to the operator due to the toxicity, flammability, or other hazardous characteristics of reagents, reaction products, or feaction conditions, and; (6) it should not result in a net increase in the amount of contamination present in and on the surface tested and should not in any other way affect the suitability of that surface for any projected future use.

The overall finding of this evaluation based on the available data was that there is no single spot test or combination of a few such tests which satisfies all of these criteria. More significantly, most available spot tests typically fail totally to meet one or more of these criteria. It is possible that modifications could be made to available tests to improve their performance in this regard. However, the developmental effort required for that purpose may be substantial, and it is not clear that improvement in the performance of a given spot test with respect to one criterion would be accompanied by similar improvement in other areas.

In summary, our view is that available spot tests do not satisfy minimum criteris for qualitative analysis of explosives/explosive residues on building material surfaces. We do not recommend that developmental effort be expended on spot test methods except for those analytes or conditions for which no more- or equally-promising methods can be identified and developed.

Notes for Colorimetric Spot Test Tables

Literature numbers:

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- 1 through 47 correspond to journal articles 1 through 47 listed in Table II-1.
- F₁ is Feigl, F. and Anger, V. <u>Spot Tests in Inorganic Analysis</u>, Elsevier Publishing Co., Amsterdam (1972) (6th English Edition).
- Feigl, F. and Anger, V. Spot Tests in Organic Analysis, Elsevier Publishing Co., Amsterdam (1966) (7th English Edition).
- (a) is Christensen, H.E. and Luginbyhl, T.T. (eds) Toxic Substances
 List: 1974 Edition, U.S. Department of Health, Education and
 Welfare, Public Health Service, Center for Disease Control,
 National Institute for Occupational Health and Safety,
 Rockville, Maryland, June (1974).
- (b) is Sax, I.N. <u>Dangerous Properties of Industrial Materials</u>, Reinhold Book Corporation, New York (1968).
- (c) is International Technical Information Institute (ITI)
 Toxic and Hazardous: Industrial Chemicals Safety Manual,
 Tokyo (1976).

Abbreviations:

cat - cat

C

dog - dog

gpg - guinea pig

hmn - human

inh - inhalation

ipr - intraperitoneal

ivn - intravenous

LD50 - lowest dose 50% kill

LDLo - lowest published lethal dose LC50 - lethal concentration 50% kill

LDLo - lowest published lethal concentration

mky - monkey

mus - mouse

orl - oral

rat - rat

rbt - rabbit

scu - subcutaneous

TVL - threshhold limit value; the concentration of an airborne constituent to which workers may be exposed repeatedly without adverse effects.

TABLE 11-5. SPOT TEST METHODS FOR ORGANIC ANALYTES

1714	Roagont	Sulvent	ž ž	Color	Threshhold	Interferences	Technique and Sampling
Cyclotel- methylene triniteanne (anc)	T1,7±0.1	Ethanot Sulfaric acid	2 E	Cold + red Strong heat + yellow/brown haderate heat + violet Add ethanol + Blue		IMX (leat vill separate); Any sugare and eldelydes; 4 out of 24 tested ultramines	Reating coquired; test tube
	J-w-14 (0.4Z)	95% ethanol; Mala feu secunds dater	z z	Tellow-red first; then with ethanol + blue + blue/ gren +		Nitrates (gliern	Tipe companies
	Brec. 3c		*	Ormpe to red		Browlde Mitrocellulous Chlorate Mitrostratch Hitrate Tetryl Nitrite PETM	Spart plate
	Cricsa		*	7:00		Bromide Chlorate Mitrate Mitroglyceria Tetryi Mitrocellulume Mitrostarch	Special property of the specia
	Fracilia - H N°- dimethyl-l- naphtbyl-mioe	50/50 acetic atd/E ₂ 0 10 gm zimc in 100 ml benerae	3	Red/red-wiolot 1 - 530/535 1 - 42,330/45,477	Commet laws 40 pg: news117 4 pg or 0.4 pg	Gluc on paper tweelupes; PETM, ultroglyceria; altrites	th coul (ptw.) given the color - threshold 40.0 mg; on japors - 0.4 mg
	3	60 ml concretrated Sulfaric acid	~	Tellow		ST Crates altractes altractes	Sport Plate
2,4 dlafter-	ĭ	Nut alcoluni IUX CaCl Nuj[Fe(CR)5NNj]	_~	Violet			Menting required; test tuber
	# PE UM	N mil	~	Ţ.		m-dimitto cumpomads	:

TASLE 11-5. SPOT TEST METHODS FOR ORGANIC ANALYTES (CONTINUED)

Testlas I ques auxiliantes Sampel I oug	Shot plate	Matter spart plate; samples taken from sail, confete and rubble	Test 10bc	Micro test lube	Micro test take	Spot pl	Mero test tube	Spot plate	Test tabe	Spot plate
Interferences	pare in standard emplosive residue	Mar Letracrac FETH RIM MMX	1, 3, 5 trinitrobenzese 2, 4 dinitrophenal attro composada	1MT 2.6 DMT trinitrobeazone other aromatic polymitro compounds	TMT 2.4 DMT refull robenzene other aromatic polymitro compounds	large maker of organics phenol resurcinol analine analine	Many arountic compounds	Various autue bases analite bases analitue peperidiue	distkylamines acatkylamines Various secondary amines	Vavtous analgons compounds
Threshabld	¥ ~						, 25.		. 0.	0.1.
Color	Blue (charges over time)	ş. 2.	Wiolet	702	25	Blue to blue/green	Blue	Yellow	# l rc	Blue
	~	318	2	2 2	2	2	2	22	F.2	2.
Solvent	Acetone-alcubul	Ethyl alcohol	Hot alcolos 10% CaCl, Naj[Fe(CR) ₅ NII ₃]	.5% multanic acid IN alkali alcohol acetic acid	.5% sutfante acid ly alkali alcolod accie acid	Concentrated amonto	Concentrated BCI ether 12 tetrahase	Ether	Ett.sr methonolite browne	
Keagent	252 aquecus (etfamiliylam moilum hydraulde	Saturated partessium hydroxide	50 mg zine dust	0.32 t-uaph-	.32 1-naph- thytamine	SZ phos- phomolybdic netd	Keloj	2,4-dtuftro- chlosobenzene	Third pitenyl-	n. 052 p. n. l. t. t. saphenul
Analyte	2,4-1MT				2.6 INT	Diplomel-			hur D. I	inle !

TABLE II-5. SPOT TEST METHODS FOR ORGANIC ANALYTES (CONTINUED)

Technique and Sampling	Spot plate		White spot plate (samples taken from soll, con-	Tested on papers, wad, cloth, leather	Wood ' tan color; detection Hait 4,0 pg	Wood • Can color; deterion limit 4.0 mg	Test tube	-	Mero cruethte
Interferencea		Aldehyde Thlo compounds Alkali sulfides Organic solvents (acctoue, methanul, ethanol)	Not tetracer: PERN RDX IBBX	Polynteo aromatic compounds			2,4 DNY 2,4 dinterophenol 1,3,5 triniterobenzene Other nites compounds		Compounds containing reactive CD ₂ and MI ₂ groups
Threshindd	8a 1			0.4 µg on paper	0.4 µg on paper	0.4 ng on paper			
Color	Turk red (changes	Red	Deep red	Red-red/orange λ = 500/503 ε = 32,270/72,272	Red) = 511/512 c = 10,445/22,272	Red-orange \(\lambda = 505/505\) \(\varepsilon = 28,510/22,272\)	Violet	Red	Brown-red
LJC.	2	34 6 3	31	47	۲5	<i>t</i> 5	* c	F. 2	2
Solvent	Alcoho!	Absolute alcohol	Ethyl alcohol	Ethano l	Et hano l	Ethanol	Hot alcohol 10% CaCl ₂ Na ₃ [Fe (CN) ₅ NH ₃]	.52 suffante neld IN alkalt alcohol acette actd	50% alcobal difute NaOH
Reagent	25% aqueous Tetramethylam- montum	Nessler's reagent	Saturated potassion hydroxide	1,3 diphenytanetre	Cyclopentanone El ₄ -NOH	Ni tromethane Et ₄ -NOI	Zine dust (50 mg)	. 3% Lenaph- thy landine	1, 2-naphtho- gofnone -4- soffonate
Analyte	TNT		-					-	

Violet	7,2	10th will benzene Hot alcohol 10Z CiCl2	10g zinc dust 10t =1 benzene 10t atcobal 10Z CiCl ₂
	ີ ວ ລ	F ₂	nit, i

TABLE II-5. SPOT TEST METHODS FOR ORGANIC ANALYTES (CONTINUED)

Technique and Sampling	sunds, Sample off cotton swales in ether; scanned, sample removed from robble; while spot plate	ec Spot	llulose	Mtrite Extraction with accione; Spot plate	ite 	Pods	(h) wood only 4.0 pg detection;
Interferences	Aldehydes, this companies, alkali suffides, prganie solvents (acetone, methanol, ethanol)	Browine Chlorate lodine Nitrate Nitrite Nitrocel Mitrostarch PETN Tetryl RBX	Chlorate Mitrate Nitrite Mitroce Nitrostarch PETN Tetry?	Nitrate Pitrocellulose RBX PETN Teiryl	Browlne Chlor lodine Nitra Nitracellulose PETN Nitrostarch RDX Tetryl	Charate Indine Nitrite Ferchlorate	Nicrite
Threshhold	2 = 5						0.4 pg usually
Color	Yellov + orange/ brown ammonia + black	Or ange - red	Slue-blue/black	Pink to red	Orange -brown	light, dirty-white precipitate	Red-red/violet 2 = 530/535
	. s 6	*	*	*	*	34	\$
Solvent	Absolute attoluil sumonts	Acetoue	Acet ove:	Actone	Acetouc		50/50 acetic acidiligo Zim dust (10 B)
Reagent		Frac (see	Diplemylanine	Gress)-ac-ld	Nitron	Procedue n,n' diethyl-l-
Analyte	Ş						

TABLE 11-6. REAGENTS FOR ORGANIC ANALYTES

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			*****	Prac	Practicability			Rigard	
Reagont	Solvent	Analyte	1.1€. ₩0.	Destruct iveness	Application	Staplicity	Toxicity	Florenchillts y	hinger
B.:tazenec			~				Suspected carelingues; 19, 4033 rec) 25 ppm in air 80 mg/m in air (b) 71.V = 10 ppm (b) (c)	Rigorans react fon	Face shield; breathing ap- paratus; Sately gogglen (c)
Bruche S g broche sulfate	100 ml concentrated sulfurle	757 767 767 767 767	ž	Acid chare organics	·		litgh (b) Inhalation highly toxic; fpr-rut 1D ₅₀ =77 mg Ivn-rbt Litox30 mg (c)	theat toxic fumes (b)	Kubber gloves; Work in draft protective clothing; Respirator; (c)
Chlorant 1 (tetrachloro- Lenzoquinone)	Ether KC10 ₃ HC10 ₃	Of pleeny 1-	2	Redux reaction		Requirer a.r dryleg, warming.	See <u>benzintulnum:</u> 1.050 = 7510 <u>14</u> 1.050 = 7510 kg	lkat + toxic tumes (b)	
Cyc lopentanone Et ₄ on	Etharot	Ē	7				11150 = 4490 #8 1.111.0 = 102 mg/kg (a) Cycloparaffilms: deadly marcotic (b)	Maderate Clammable (b)	:
6-dfaltro-	Ether	Diplenyl-	2 م	Forms quinoldal zultterion condensation primicts				18 (a) 19 H	Haderate explosive when exposed to flame (b) slack or heat expludes (c)
1, 3 Diptemyl- acctone Etginst	Ethan	Ē	£\$			Commercially avidable		; ; ; ; ;	
	2 2 2 2	PETA	=	Acid chars organics		18 also an analyte	~200 mg kg 102 mg/kg	:	Explositve

TABLE II-6. REAGENTS FOR ORGANIC ANALYTES (CONTINUED)

.

					Pract Icability			Bazard	:
Reagent	Solvent	Analyte	7. I.f.	lest ruct i venuss	Application	Stoplicity	Toxicity	Florenchilley	Danger
Phosphomolyhelle acid (52)	Anmonta	Diphenyl- asine		redux reaction					
Por as loss cyan's de	ZN IICI (later)	2,4 d1- nitro- phenot	2	Presumed product: phenyllydoxy- lamines		Requires bearing	Skin irritunt taxic (b)	Roderate flammable, cults hydrocyanic Acid on heating: highly toxic gas	
Potastom trydroxfde	Ethyl	THT 2,4-IMT	31				orl-rat LD50=365 kg (c) highly toxic for subalation (b)	reacts with water or steam to pro- duce consite solution - toxie (b)	Corrosive Hquid (b) rubber gloves face saled
Fractine (, 35 gm) N,N'-dimethyl-1- nophthylomine (, 35 gm)	\$0-\$0 acette 3c1d - H ₂ 0 10 gm Zn dust henzene (100 ml)	PETN NG RDX	£3				14150=500 mg/kg 14150 ~1400 mg/kg (a)	Hoderate heat >	Explosive under shock or heat (b)
(b) tool tractu (J mg) (1, 2, 5, 8-tetra- hydroxyanthra- qoloone)	40 mt concentrated sulfuric	RDX	F ₂	Nitric acid splits off	Stir 20 sec; wait 20 min. preciscly				
25% aquions tetramethy- lamman un hydroxfde	acetone	TNT m-d1- nitro- benzene 2-4 DNT	14			Clear mobile Hquid	Toxic - powerful caustir (b)		Corrosive Huni
Thymest	Suffurte actd Ethanol	KIJX	_	Acid chars organic		Requires heating to 190°C	S14ghr toxicity (b) ori rat th\$0=900 aB ori rat th\$0=1800 aB ori rak th\$0=1800 aB vri rak th\$0=800 aB	Toxic funcs upon heating (b) Finnantic (c)	Rather gloves lace shield gas mask (c)
	T								

TABLE II-6. REAGENTS FOR ORGANIC ANALYTES (CONTINUED)

	J.rilem _Q	Heathly arlac analysis meressary (c)				Explosive store anay from beat; relater gloves, respirator, plastic clothes (c)		Highly captonive chas cr-
Hazard	Flammability	Slight heat ' foxfe fum-15 (b.)			Heated + toxic lodine functi (b)			
	Toxicity	teagent lasts 1D ₅₀ v2 g/kg. 2 months if refrigerated 1Dlo=150 mg/kg. (a) Highly toxic, bladder cancer (b) orl-rat 1D50=79 wg scu-mus TDLO=25 kg scu-dog TDLO=400 mg (c)			KI: Prolonged absorp- tion +lodism, skin rash (b)	1.050 = 940 mg/kg 1.01.0 = 102 mg/kg (a) (a) (b) 111 ghly toxic Try (ACCHI rec) 100 ppm in air 250 ug/m³ in air (b) (in) (in) (c)		May resemble p-nitrophol; If so, high toxicity (h)
	Simplicity	Reagent lasts 1150 v2 g/kg 2 months 1f refrigerated 1100=150 mg/l Highly toxic, bladder conce orl-rat 11550 scu-mus TD1,0 scu-dog TD1,0						
Practicability	Application							
Prac	Destructiveness	Ar - N0 goes to Ar - OK	Acid chars organics	Forms: indophenol dye				Oxidized to merlquinoidal compounds
:	٠. او.	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	36	₂	4.3	15	34	F.M
	Anaiyte	2,4 d1- nitro- phenol; TWT; 2,4 DWT; 2,6 UWT; 1,3,5 tr1- nitro- benzene; PETN; RDX	RDX	TNT	NC TNF	I X I	X:	dfphenyl- anine
	Solvent	0.5% sulfante acid 1N alkall alcohol acet le acid	95% ethanol	5 0% alcohol dilute NaOH	Alcohol	Rthano 1	88% formic actu	161
	Rengent	Griess reagent (c-naphthylawine) (l-naphthylawine)	J-actd (6-amino-1- naphthof suffonte acid)	1, 2_naphtho- qu Inone- 4-su I fonate	Nessier's Keagent (R1) (1861 ₂) (K011)	Hitromethane	Nitron	p-ntrosophenol

TABLE 11-7. SPOT TEST METHODS FOR INORGANIC ANALYTES

THE STATE OF THE PARTY OF THE P

Technique and	Filter paper ed	Spot plate	Spot plate	Heated in boiling water bath for 5 minutes Spot plate	Filter pape	Spot plate		Micro lest tube todiling pater	
Interferences	Metal tons and cut2, Mg, Sn, Sb if reagent treated with ammonia, only Ag and Tl interfer			Cu, Co, NI Not Cr ⁴⁶	Not Group 111 Metals	си ⁺² , со ⁺² , и1 ⁺² гс ⁺³		Netals Allete	
Threshlold	٠.05	۲.80.	.05y	Microgram	. 06 y	.25 _Y	.25م		
Culor	Red	Brown green/blue	8 LC	Reddish viol°C	Orange-yellow	Vlolet	91116	Violet	Plak
No.	يـــ بــــ	1	<u>.</u>	:				- i	38
Solvent		Formaldehyde 10% NaOH 10% KCM Alcohol	Piperidine alcohol 20% sodium thiosulfate 20% sodium tartrate Suturated sodium fluoride		lleated 2N 11 ₂ 50 ₄	Concentrated 12504 202 Sulfosal-trylle acid	Moderate concentration audium per- oxide Aeetie acid	Alcohol 0.5N caustic alkati	25% Acette acto
Reakent	Fc(u,n'-d1p ₁)1 ₂	OI-p-n(tropheny)- curbaz tde	Glyoxal bis- (2-nydroxyan11)	Chenta ac14	Acid Altzaria KC	Alkall bypobro-	Beuz l'Afne	o-dlotrobensene	Seal from
Analyte	3	!	3	. c	- G- J-	9 ₄ v	÷ č	Ē	£

TABLE II-8. REAGENTS FOR INORGANIC ANALYTES

				Prac	cicability		<u> </u>	#483FG	
Reagang I	Solvent	Analyte	Mo.	Destructiveness	Application	Simplicity	Toxicity	Flatmability	Danger
eid Alizaria	TH H ² RO. ²	Ce ⁻³	F _L		Cannot be reshed out of paper vith H ₂ D or icids	Requires			1
Lind Commerce	Toncencrated 4.0 50; 107 sulforate icytic acid	ce ⁻⁶	7,	Forme CcO2	 				!
Pensidine	Sodium peroxide acetic hold	Cr ⁻⁴	7,	Omidized to chromate			-Any exposure extremely has- ardous; poison label (b) -Carcinogen orl-dom LDLo 100 mg/kg (c)	-Heat - uecomposition hazardous fuces (b)	-'ubbat gloves:
ienta 401d .2-dimino- cyclonexane- i,d.N'.31- tectascetic .ctd)		:e+3	F.			Acquires boiling water bath			
- iinitrobenza	Alconol 9.5% causcic gikali	Po	F ₁	Forme eikeli seld of quinoidel nitrol-mitro acto	i		m-inicrocensene: TLV = 1 mm/mg orl-cac iblo = 27 mm/kg (c)	<u>n-dinitroberzene</u> : Blight	m-distrobenze High explosive hazard (b) Special outdoo storage advise (c) Rubber gloves; breathing 19- Aratus (c)
-p-Hittonnenvil	102 NacM 102 KCN 40% formeldening	Cd Cd	r ₁	Forms (d(OR)		<u> </u>			PARECUS (C)
(37 '-410) 312 -410 YELD	Amonia	C d	F ₁	Forms Cdt,			For so'dipyridyl see pyridine (b) 2vridine: ITV = 5 ppm in aie (b)	Pyridine: hast = decomposi- tion, hexardous fumes, highly flamable (b)	Pyridine: explosive vapor special scorage (c) Tubher glowes(c) Feathing apparent
(2-nvdroxyanil)		લ	£.	forms a colored chelsts	1 	Reduires besds of resin	olymeat: moderate (5) orl-rat LDLo = 100 mg/kg (c)	Sluckal: slight (5)	Clyonal: rubber gloves breaching in- partus (c)
iedium	Aceste Acid	75	18	1	i		;		

D. FINDINGS AND CONCLUSIONS:

Among the overall findings of the literature search and spot test review are the following:

- The literature search indicates that most of the prior work in this area has been done by law enforcement agencies, and those agencies have, for the most part, concentrated on the application of spot test methods for the detection of explosives: quantitative determination of explosives has not been a major concern, except to the extent that these investigators have attempted to demonstrate the range of concentrations to which their qualitative detection methods could be applied. The Army, also, has relied mainly on spot test methods up to now in explosives contamination assessment work. However, as noted above, available spot tests do not satisfy minimum criteria for qualitative analysis of explosives/explosive residues on building material surfaces.
- Other investigators have reported explosives determination techniques involving analytical methods ranging in sophistication from visual examination of residue under low power magnification to mass spectrometry, including in between almost all readily available modern analytical techniques. Sampling for quantitative analysis remains a weak spot. Thus, different methods for qualitative and quantitative analysis may be required to satisfy the Army's requirements.
- The use of a qualitative survey obviously must not preclude the subsequent use of a quantitative method if two different methods are used for qualitative and quantitative analysis, application of the qualitative method should not result in the destruction or loss of analyte. Many of the methods described in articles collected in the literature review result in loss or destruction of analyte and thus may not be applicable to the present problem.
- Recommended sampling and analytical methods should not result in a net increase in the amount of contamination through the application of toxic or hazardous reagents to the surface to be tesced, nor should they present any other hazard to the operator. Many of the materials described in the literature review require the use of toxic reagents and thus for this reason also may not be applicable to the present problem.
- Surface contamination of the type the Army is concerned with is most likely to have resulted from spillage or dusting of solids or from spillage of process liquors or liquid wastes. The latter situation presents the problem of sampling compounds which may have penetrated into porous surfaces such as wood or concrete. Most existing methods do not address this issue.

E. SAMPLING PROTOCOL SELECTION:

Based on the review of the available literature and discussions with the Technical Project Officer and other persons having expertise in trace analysis and explosives technology, the following sampling protocols were identified as candidates for developmental testing:

- continuous monitoring of organic analyte vapors using a portable gas chromatograph;
- 2. evaluation of existing U.S. Army equipment developed for the vapor phase detection of CW agents;
- in-situ formation of charge-transfer complexes with visual identification;
- 4. UV irradiation of suspected contaminated surfaces with subsequent detection based on thermal imaging or UV photography of the irradiated surface;
- 5. Solvent extraction using alternative procedures to conventional wipe or swab methods.

Protocols 1, 2, 3, and 4 were candidates for qualitative detection of explosives/explosives residues; protocol 5, compined with modified versions of existing USATHAMA methods for the determination of explosives was intended for quantitative determination. Detailed discussions of the developmental testing of these qualitative and quantitative methods are presented in the respective sections of this report.

·III. QUALITATIVE METHODS DEVELOPMENT

A. INTRODUCTION:

The qualitative methods selected for developmental testing included:

- 1. continuous monitoring of organic analyte vapors using a portable gas chromatograph;
- 2. evaluation of existing U.S. Army equipment developed for the vapor phase detection of CW agents;
- 3. in-situ formation of change-transfer complexes with visual identification;
- 4. UV irradiation of suspected contaminated surfaces with subsequent detection based on thermal imaging or UV photography.

Detailed discussions of the developmental testing of these methods are presented in the respective sections below.

The objective of this testing was development of procedures for the rapid qualitative determination with 90% confidence of the presence/absence of the compounds of interest lown to a level of 5 $\mu g/10~cm^2$ in a given building. The approach used to achieve this objective involved in each case the spiking with known amounts of analytes of new, uncontaminated samples of each of the surface types of interest obtained from building materials dealers. Samples of conductive non-sparking flooring were not available for this purpose.

Practical determination of the confidence interval associated with a qualitative analysis performed in the field would have required access to a building where the nature and extent of contamination was known. This condition could not be satisfied by those AAP's to which access was obtained. Therefore, detection limits for positive compound identification are reported.

At the direction of the Technical Project Officer, emphasis was placed on the development of procedures for organic analytes. No methods for inorganic species which would represent substantive improvement over existing spot test methods were finally identified.

B. ANALYTE DETECTION USING CONTINUOUS VAPOR PHASE MONITORING:

The detection of explosive vapors using gas chromatography and other analytical techniques has been the object of considerable investigative effort. In fact, several of the commercially-available explosives detectors are based on this principle. Available explosives vapor detection methods have been used with varying degrees of success for the detection of bulk explosives as, for example, in airport and aircraft security operations.

Detection of explosives/explosive residues on building materials surfaces using continuous vapor phase monitoring could offer the following advantages:

- By means of one or a few measurements, all the surfaces composing an entire enclosed area could be effectively screened for the presence of explosives;
- The analyses could be performed on or close to a real-time basis, making it possible to conduct "walk-through" surveys.

Unfortunately, however, there are several potential difficulties with using continuous vapor phase monitoring for detection of explosives on surfaces, including the following:

- The concentrations of analytes on surfaces which are of interest in this study are very low;
- Further, the vapor pressures of most of the analytes of interest are very low under ambient conditions;
- Most of the analytes of interest are strongly polar and thus adsorb strongly on surfaces with which they come in contact.

These factors have, in fact, largely precluded the effective use of vapor phase detection methods for many applications where the detection of other than bulk quantities of explosives was attempted.

Two types of recently-developed analytical instruments were evaluated in this study to determine whether their operating and performance characteristics made possible the detection of vapors derived from explosives in air immediately adjacent to surfaces spiked with explosives/explosive residues. One was the Photovac 10A10 Portable Gas Chromatograph (Photovac, Inc., Thornhill, Ontario, Canada), a portable gas chromatograph with a photoionization detector for which detection levels down to parts per billion for various compounds including nitro-compounds are claimed. A technical representative of Photovac, Inc. visited Arthur D. Little, Inc. laboratories to discuss the application of the Photovac 10AlO Portable Gas Chromatograph to continuous monitoring of organic explosive vapors and to demonstrate the performance of that instrument used in its continuous monitoring mode for sampling of air immediately above a surface spiked with known amounds of selected explosives. During that demonstration, the Photovac 10A10 failed to give any observable signal when used to sample the air immediately above the bottoms of Pyrex glass beakers spiked with the equivalent of 125 µg/cm² of RDX, 2,4-DNT, and DPA.

The Arthur D. Little, Inc. Project Manager and the Technical Project Officer also visited U.S. Army C.S.L. Laboratories to view and assess the performance of U.S. Army CW agent field detection equipment when used to sample explosives vapors. Three ionization detectors, including the model in current use and two prototype instruments, were evaluated by sampling the air immediately above the bottoms of Pyrex glass beakers spiked with various concentrations of RDX, 2,4-DNT, and DPA. Each of the detectors gave what appeared to be an observable response at or near the agreed-upon detection

limit of $0.5~\mu g/cm^2$. However, in no case was that response sufficiently large to assure unambiguous detection. It was judged that none of these instruments appears at this time to possess sufficient sensitivity to warrant further investigation in this project.

C. ANALYTE DETECTION USING FORMATION OF CHARGE TRANFER COMPLEXES WITH VISUAL IDENTIFICATION:

1. Background.

Charge-transfer complexes, particularly molecular addition .com particularly molecular addition .com have been used for years in the isolation, purification and idea of organic compounds. Among the better known examples are the compounds of 2,4,6-trinitrophenol (picric acid), 1,3,5-tr:..trobenzary and 2,4,7-trinitrofluorenone. These nitroaromatic compounds are chargetransfer "electron-acceptors." Their complexes with "electron-dor rs' are usually formed in 1:1 ratios of acceptor:donor, and the covexhibits properties differing from the individual components, e.g., color, crystal structure, melting point, etc. The bond a rength of the addition compounds varies from very weak--on the order of van der waal's forces -- to moderate strength such as in hydrogen bonding. The wimple. aromatic hydrocarbons, e.g., durene, naphthalene, and anthracer, are typical donor compounds whose complexes have the weaker type of bonds such that the solid complexes can be readily dissociated by log a of the donor through volatilization at or only slightly above room temperature. Solvent action can also be used to remove the donor compound from the complex.

In the interaction of nitroaromatic hydrocarbons with the simpler aromatic materials, visible color due to complex formation would be the simplest method of detection. In the absence of a good color contrast, the known property of nitrogromatics to quench the fluorescence of the other aromatic compounds could be used as the basis for a less simple detection scheme. Of the donor compounds mentioned above, however, only anthracene has a fluorescence at visible wave lengths such that it could be used without an instrumental detector. Still another combination of a physico-chemical interaction of a fluorescent donor with the quenching nitroaromatic acceptor compound as part of an energy-transfer system might also be used to increase the sensitivity of detection. Work performed by Arthur D. Little, Inc. on detection of polycyclic aromatic hydrocarbons indicates that less than nanogram quantities of anthracene can be detected by virtue of that compound's energized fluorescence. The interference of nitroaromatic compounds with such a test should thus permit their detection at a corresponding level (cf. Interagency Energy/Environmental Report EPA 600/7-78-182, September 1978).

In summary, in-situ formation of charge-transfer complexes for the detection of explosives on building materials surfaces offers the following advantages:

- (1) Sensitivity down to the desired detection limit of $0.5 \mu g$ of analyte/cm²;
- (2) Speed: the chemical reactions used in this application occur under ambient conditions; relatively easy and straightforward methods for dispersal of the required reagents over large areas of the surface types of interest are available; and visual observation of colors or quenching is used as the detection method;
- (3) Manageable hazard during and subsequent to use as compared to similar spot test methods; and
- (4) Reversibility: the charge-transfer complexes used in this application can be destroyed relatively easily leaving the original analyte intact and available for subsequent quantitative testing.

2. Preliminary Experiments

Whatman No. 42 filter paper was used as a substrate in place of samples of the actual surface types of interest in initial spiking experiments. Filter paper or the equivalent is customarily used in this type of work since it provides a convenient means for manipulating the small amounts of chemicals involved and also provides a nearly ideal visual background for visualizing the colors or fluorescence of reaction products.

Acetonitrile solutions of three of the organic analytes--2,4,6-TNT, 2,4-DNT, and RDX--were applied to Whatman No. 42 filter paper in quantities sufficient to yield concentrations equivalent to 0.5 μg and 50 μg of analyte/cm² (= lx and 10x the agreed-upon detection limit of 0.5 μg /cm²). The acetonitrile was allowed to evaporate, and the spiked filter paper was then treated with cotton swabs which had been immersed in acetonitrile solutions containing 100 μg /mL of the electron-donor compounds durene, hexamethylbenzene, naphthalene, and anthracene.

The filter paper was then examined for (1) the presence of colored reaction products, or (2) quenching of reagent fluorescence when examined under a UV lamp. First priority was assigned to identifying a reagent which gave a colored reaction product which, at all concentrations, was clearly visible to the unaided eye under ambient lighting conditions and was clearly distinguishable from any potentially interfering colors or surface irregularities. Detection of a reaction by observation of quenching of reagent fluorescence was considered acceptable only if a colored reaction product could be identified; in that case, the observed quenching should satisfy the same criteria as above.

In an effort to identify an electron-donor compound which would yield a colored reaction product with these analytes, the compounds N,N-dimethylaniline and diphenylamine were also evaluated using the procedures described above. While diphenylamine is also an analyte in this program, it was nevertheless considered useful to evaluate its electron-donor properties since it is often cited in the literature as a strong electron-donor and could thus serve as a useful model compound.

At the 50 $\mu g/cm^2$ level, 2,4,6-TNT formed a reddish-orange spot with both N,N-dimethylaniline and diphenylamine; 2,4-DNT formed a pale yellow spot with diphenylamine only; and RDX failed to form an observable colored spot with either compound. At the 0.5 $\mu g/cm^2$ level, none of the analytes formed an observable colored spot with either compound. However, when spiked filter paper treated with anthracene was observed under ultraviolet illumination (254 nm), quenching of the anthracene fluorescence was observed for all analytes at both concentrations.

On the basis of these initial positive findings, additional experiments were performed on the nine analytes listed below:

Analytes Tested

NG Nitroglycerin PETN Pentaerythritetetranitrate RDX Cyclotrimethylenetrinitramine TNT 2,4,6-trinitrotoluene TNB 1,3,5-trinitrobenzene 2,4-DNT 2,4-dinitrotoluene 2,6-DNT 2,6-dinitrotoluene DNP 2,4-dinitrophenol Tetryl 2,4,6-trinitrophenylmethylnitramine

Each analyte was spiked on Whatman No. 42 paper at levels corresponding to 0.5, 1.0 and 100X the specified detection limit (0.5 $\mu g/cm^2$). As in the preceding experiments, cotton swabs which had been immersed in an acetonitrile solution containing 100 $\mu g/mL$ of anthracene was gently drawn across the filter paper. After the acetonitrile had evaporated, the paper was examined under 254 nm UV illumination.

All of the analytes except NG and PETN were detected at the 100% level without application of anthracene; the detections were as dark spots on the light paper background. (In visible white light, the DNP and Tetryl spots were yellow.) When treated with anthracene, all of the analyte spots were seen as dark spots on the fluorescent anthracene background.

At the 1X $(0.5~\mu g/cm^2)$ level, again only NG and PETN spots were not evident until treated with anthracene. Even at this level, they did show a weak quench of the anthracene fluorescence. The other compounds, evident without anthracene, showed increased contrast (dark/white) with the reagent.

At the 1/2X (0.25 $\mu g/cm^2$) level, none of the materials were detected until treated with anthracene. PETN, NG, and RDX detections are questionable at that level. Taken together, these findings suggested that detection of the three analytes tested at the agreed-upon detection limit of 0.5 $\mu g/cm^2$ was well within the capabilities of the technique.

3. Formation of Charge-Transfer Complexes Directly on Surfaces.

In an additional set of experiments, analytes were spiked directly on clean samples of the surface types of interest. The acetonitrile/anthracene solution was sprayed on the spiked surface sample using a spray bottle. In each case in which the quenching of the anthracene was positively identified, a lower concentration of the analyte was spiked on a clean sample of the surface type being examined on the experiment repeated. The lowest concentration at which quenching of the anthracene fluorescence could be positively identified was taken as the detection limit for that analyte-surface combination. The resulting detection limits are listed in Table III-1.

An additional finding of this work was that several spray applications (i.e., 3-4) of the anthracene/naphthalene reagent are necessary to achieve a uniform fluorescence background on brick and concrete. With a single application of reagent, surface irregularities appear as slightly darker areas and cannot be distinguished from analyte.

It was also observed in these experiments that the presence of dirt or other foreign matter on metal and wood surfaces prevented uniform spreading of the reagent over the surface. Also, care is required on metal and wood surfaces to avoid puddling or running of the reagent, which may result in removal from or dilution within the area being examined of any analyte present.

4. Solvent Lift Technique for Sampling of Surfaces.

The analyte detection limits shown in Table III-1 are higher for all analyte-surface combinations than for the same analytes on filter paper. In an effort to determine whether lower detection limits could be obtained, alternative procedures for isolating the analyte from a surface prior to treatment with the acetonitrile/anthracene solution were evaluated.

BY FORMATION OF CHARGE TRANSFER COMPLEXES DIRECTLY ON SURFACES LABORATORY ANALYTE DETECTION LIMITS (MICROGRAM/cm²) OBTAINED TABLE III-1.

Analyte			Surface		
	Hetall, 2	Concrete	Transite	Brick	Wood ²
£	200	320	453	20	240
PETN	200	320	453	90	400
RDX	200	320	144 3	160	400
TAT	• 009	320	30,	160	240
TNB	42	320,	30 %	1604	160
2,4-DNT	13	100	14	\$0 \$	160
2,6-DNT	13	100	. 14	204	96
DNP	424	100 5	30,	90	96
Tetryl	424	320 4, 5	30	20	1605

All analytes could be seen at the reported concentrations with the unaided eye as crystals on the metal surfaces. 2Nood and metal surfaces were precleaned with acetone to assure uniform spreading of reagent.

³No increase in the analyte/background contrast was observed at analyte concentrations up to 300 µg/cm².

analytes may be seen at "Reported concentrations are for positive identification: lower concentrations.

⁵DNP and Tetryl could be seen at the reported concentrations with the unaided eye as yellow stains. The best results were obtained using the following procedures: 1) Whatman No. 42 9.0 cm filter paper circles were saturated with 0.5-10 mL acetonitrile; 2) the wetted filter paper was pressed against the surface of interest; 3) the filter paper was allowed to remain in place until the acetonitrile had evaporated; 4) the filter paper was removed and stored in 100×15 mm disposable plastic Petri dishes (Fisher Scienctific Co. Cat. No. 8-757-12) until analyzed.

Using these procedures, the detection limits are shown in Table III-2 were obtained. Comparison of these results to those in Table III-1 indicates that the detection limits obtained using the solvent lift technique are lower in almost all cases than those obtained by formation and detection of charge-transfer complexes directly on surfaces. The solvent/lift approach also eliminates the analyte dilution and reagent puddling problems which may accompany the accidental application of excess reagent.

5. Field Evaluation.

During the week of April 19, 1982, visits were made to Holston Army Ammunition Plant in Kingsport, Tennessee and Joliet Army Ammunition Plant in Joliet, Illinois to evaluate under field conditions the charge-transfer complex formation solvent lift sampling protocol described above. A total of 195 filter paper lift samples were collected at the two installations using the procedures described above. Seventy filter paper lift samples were obtained at Holston AAP and 125 were obtained at Joliet AAP. Six locations in five different buildings at the two installations were sampled in this manner. Solid samples were also collected from four buildings at the two installations. In each case, the specific locations sampled were those which personnel familiar with present or former manufacturing processes performed at these installations suspected were contaminated with the explosives of interest. A complete inventory of all samples is included in Table III-3.

The variety of buildings and locations sampled represents most of the sampling conditions likely to be encountered in the field. It was noted, however, that even among nominally identical manufacturing lines and buildings at a given installation, substantial differences were encountered between equipment types and configurations, previous types and of intensities of usage, etc. The significance of this observation is that the development of a generalized sampling plan would probably not be possible or useful.

Due to safety restrictions and the absence of readily accessible AC power supplies, only a few filter paper lift samples were analyzed in the field for demonstration purposes. In all other respects, however, the filter paper lift approach proved to be easy and efficient to use. All samples were returned to Arthur D. Little, Inc. laboratories and were analyzed within two weeks. Several samples which were suspected to be contaminated with explosives were reanalyzed at various times over

TABLE III-2. LABORATORY ANALYTE DETECTION LIMITS (MICROGRAMS/cm2) USING SOLVENT LIFT TECHNIQUE

C

			Surface		
Analyte	<u>Metal</u>	Concrete	Transite	Brick	Mood
RDX	80	100	20	100	0.5
TXT	'n	20	90	50	0.5
2,4-DNT	\$	95.	S	50	0.5
2,6-DNT	٧	20	50	90	0.5
Terryl	ĸ	100	\$	50	0.5

TABLE III-3. INVENTORY OF SAMPLES COLLECTED AT HOLSTON AND JOLIET ARMY AMMUNITION PLANTS

Site	Building	Area S.	ample No.	Sample Type
Holston	: 1	top of wooden drain cover	1 2 3 4	filter paper lift
Holston	Il	bottom of wooden drain cover	5 6 7 8	filter paper lift
Holston	11	bottom of concrete drain basin	9 10 11	filter paper lift
Holston	Il	wall frame base (wooden)	· 12 · 13	filter paper lift
Holston	Il	metal wall frame	14 15	filter paper lift
Holston	11	base of concrete wall	16 17	filter paper lift
Holston	Il	two feet from floor on concrete wall	18	filter paper lift
Holston	11	corner of concrete	19 20	filter paper lift
Holston	11	one foot from floor on a concrete wall	21 22 23 24 25 26 27	filter paper lift
Holston	11	bottom of wooden door frame	28	filter paper lift
Holston	Il	one foot up on a concrete wall	29 30 31 32 33	filter paper lift

TABLE III-3. INVENTORY OF SAMPLES COLLECTED AT HOLSTON AND JOLIET ARMY AMMUNITION PLANTS (CONTINUED)

Site	Building	Area	Sample No.	Sample Type
Holston	11	concrete floor	34	filter paper
		(area covered	+	
		15 1/2' x 14')	55	
Holston	Il	concrete pump	56	filter paper
		base	↓	lift
			63	
Holston	11	metal strip	64	filter paper
		a long face	65	lift
		of pump base	66	
Holston	11	drain basin	67	white powder
		pump base	68S	concrete
			69S	concrete
			70S	concrete
Holston	D ₄	flooring	718	asphalte
	(RDX	wall	72S	brick
	reaction	baseboard	73S	tar material
	building)	door frame	74S	wood
	•	tank base	758	white powder material
		equipment base	76S	concrete
		pump base	77 S	concrete
		pump base	78S	concrete
		wall base	798	black tar material
		drain base	80S	black powder
Joliet	TNT	second floor	101	filter paper
	shell	melting bay	+	lift
	loading	area (concrete)	162	
Jolier	TNT	first floor	163	filter paper
	shell	loading bay	↓	lift
	loading	(concrete)	186	
Joliet	DNT	first floor	187	filter paper
	sweathouse	drain area	+	lift
		(concrete)	194	
Joliet	DNT	transite	200	filter paper
	sweathouse	wall paneling	†	lift
			204	
Joliet	INT	concrete	205	filter paper
	washhouse	floor	+	lift
			209	
				Arthur D Little

Arthur D Little Inc

TABLE III-3. INVENTORY OF SAMPLES COLLECTED AT HOLSTON AND JOLIET ARMY AMMUNITION PLANTS (CONTINUED)

Site	Building	Area	Sample No.	Sample Type
Joliet	TNT washhouse	concrete sump basin	210 + 213	filter paper lift
Joliet	Tetryl packaging	non sparking floor	216 217 218	filter paper lift
Joliet	Tetryl packaging	concrete drain basin	219 + 222	filter paper lift
Joliet	Tetryl packaging	non sparking floor	223 .	filter paper lift
Joleit	Tetryl	sheet metal dust collector	224 225	filter paper lift
Joliet	TNT unloading	second floor flooring	3015	wood floor covering
Joliet	Tetryl packaging	floor	3025	non sparking floor
Joliet	Tetryl	drain basin	3035	white powder

an additional two-week period and in each case the observed results were indistinguishable, indicating that no observable loss of analyte occurred upon storage of samples for up to one month.

The solid sample 675 collected from a floor drain in Building I-l at Holston AAP was analyzed separately by x-ray diffraction and was found to consist mainly of RDX. Acetonitrile washes of several filter paper lift samples collected from the same general area were also analyzed independently by a wet chemical method for identification of RDX (ref. Dept. of the Army Technical Manual TM 9-100-214, Military Explosives, No. 1967, p. 12-4) and were found to give positive results for that analyte. Several additional filter paper lift samples from nearby areas which contained amounts of dirt and debris similar in quantity and appearance to that on the samples described above gave no indication of contamination. Taken together, these observations suggest that certain of the areas sampled in Building I-l at Holston AAP were indeed contaminated with RDX, that the charge-transfer complex formation sampling protocol gave a positive indication of contamination in and around areas where contamination was known to exist, and, in areas further removed from the known contaminated areas, the chargetransfer sampling protocol gave fewer or no indications of contamination. The latter observation, in particular, suggests that the presence of dirt and debris of the type found throughout the building did not result in false positive findings. This is precisely the outcome which had been desired.

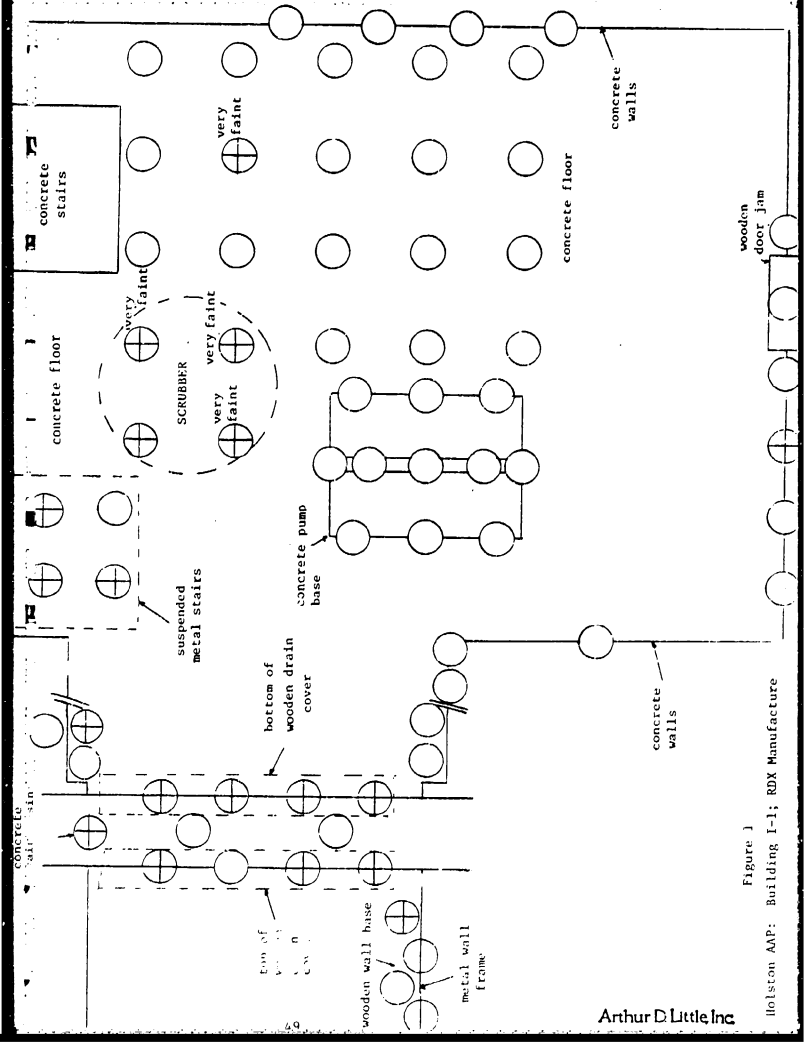
The results of analyses of all filter lift samples are represented diagramatically in Figures III-1 through III-6. In those figures, small circles indicate the locations from which filter paper lift samples were collected. Open circles O represent a finding of no apparent explosives contamination; crossed circles O represent a finding that the location sampled was contaminated with explosives. Findings which were questionable due to very faint fluorescnece are denoted by the appropriate notation next to the corresponding location in the figure.

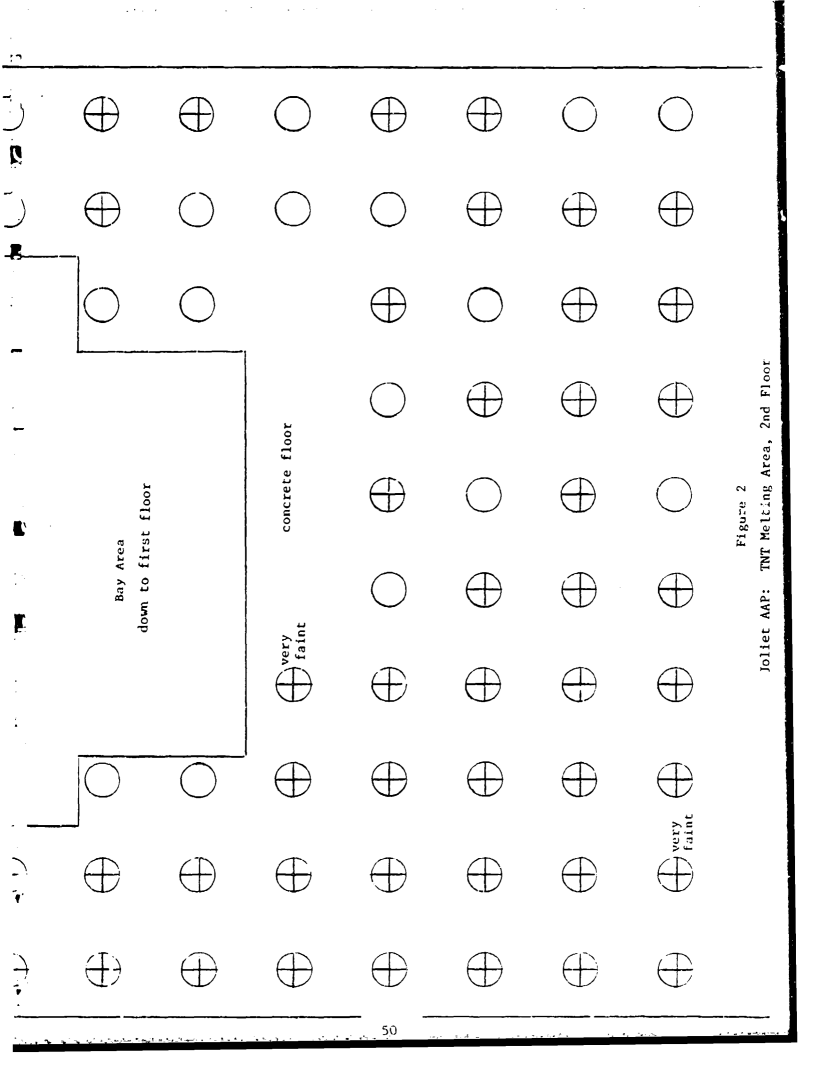
D. ANALYTE DETECTION USING UV IRRADIATION AND THERMAL IMAGING:

1. Background.

An analytical approach utilizing UV irradiation and thermal imaging technology was evaluated for its potential applicability to the detection of trace levels of explosives on building materials surfaces. The principles underlying this approach involved the following:

The area of interest is irradiated with an ultraviolet illumination source matched to the absorption characteristics of the analyte. The radiation that is absorbed must necessarily heat the material. A small change in temperature should result and thermal imaging technology might be used to sense that change in temperature, provided that diffusion of the heat into the substrate does not proceed too rapidly.





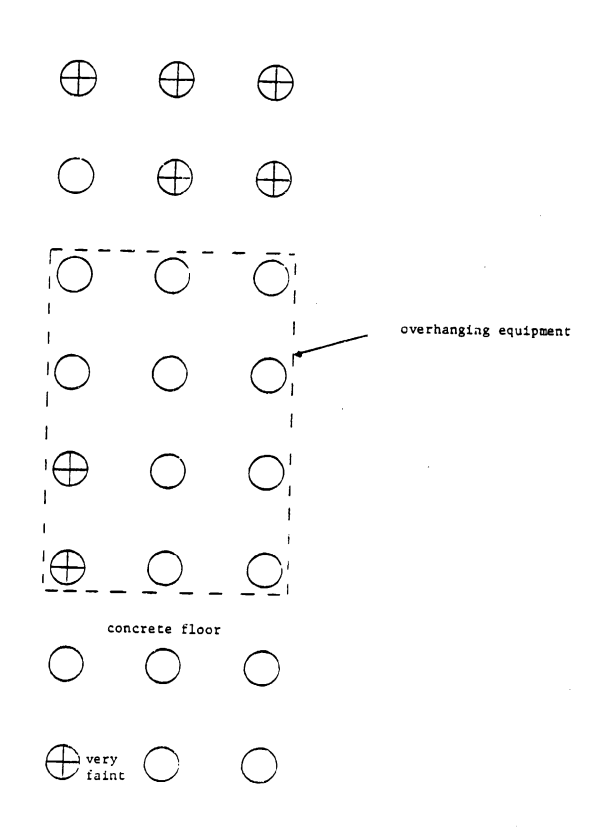
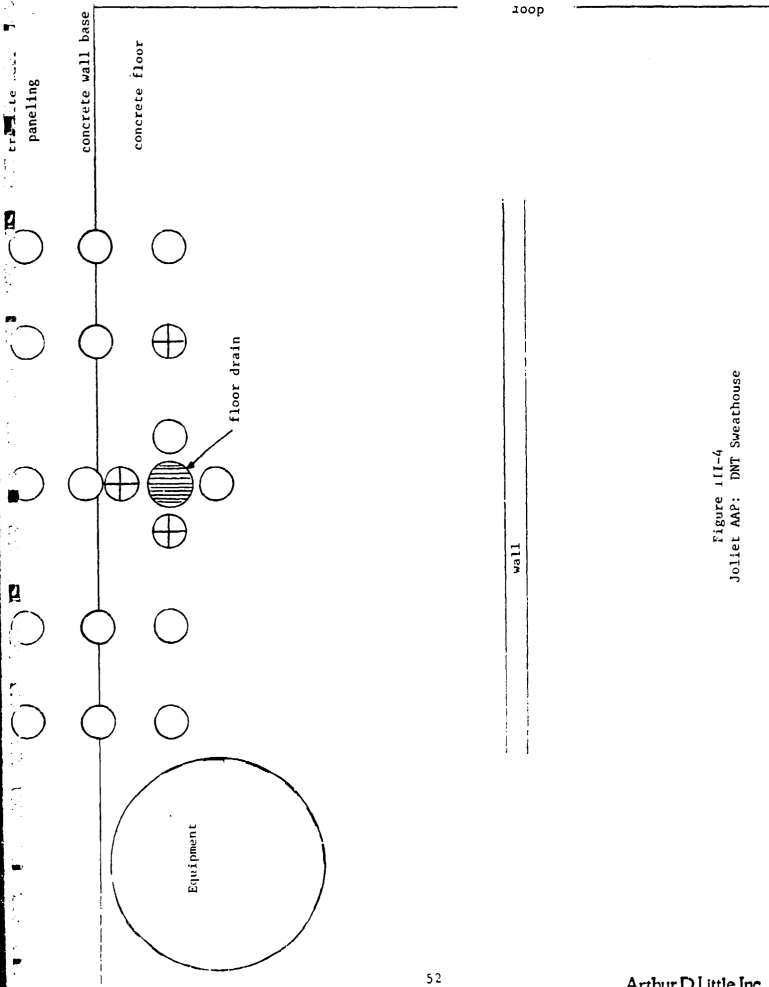


Figure III-3

Joliet AAAP: TNT Loading Area, 1st Floor



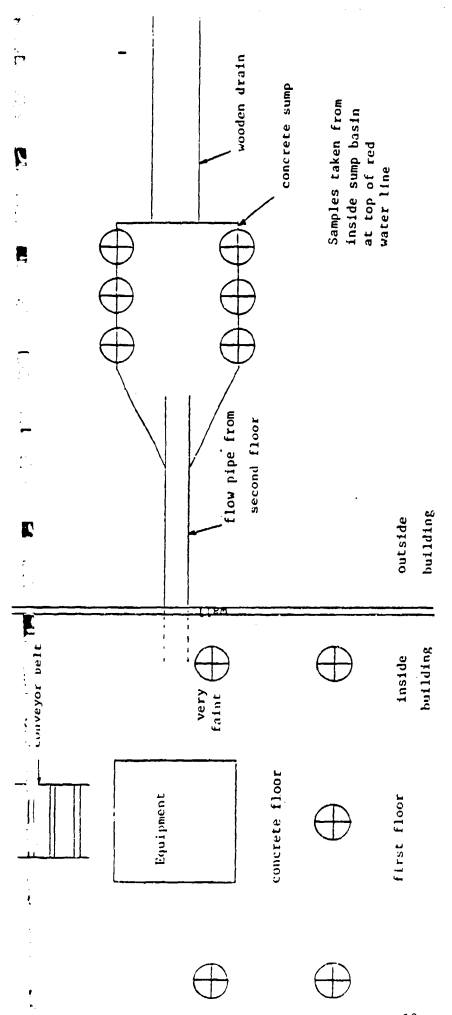


Figure III-5 Jolfet AAP: TNT Wash Building

C

7

7

E

K

Figure III-6

Joilet AAP: Tetryl Packaging Building

The technical feasibility of this approach was evaluated by a theoretical analysis which showed that the estimated temperature increases accompanying UV irradiation of trace levels of the analyses on surfaces were apparently within the measurement capabilities of commercially available thermal imaging instrumentation. Subsequent laboratory experiments demonstrated the practical applicability of the approach. The work completed in each of these steps is summarized in the respective sections below.

2. Theoretical Analysis of the Method.

a. Principle. The explosives of interest are known to have absorption bands in the middle IR and in the UV but are not fluorescent in the UV. Thus, irradiation of these compounds with UV or IR light of the proper wavelength will result in energy absorption accompanied by a rise in temperature. That temperature rise may provide the means for detecting the presence of the explosives on building materials surfaces. Because modern instrumentation is capable of precise measurement of even very small temperature changes (e.g., $\Delta T \simeq 0.1^{\circ}C$), the method may be extended to the detection of trace levels of explosives. Discrimination of analyte signal from background temperature fluctuations can, in theory, be achieved by careful selection of the incident radiation.

A detection method based on this principle would involve illuminating the wall with UV irradiation of specific wavelengths tuned to the suspected material's absorption bands and sensing the temperature rise due to the absorbed radiation. The absorption coefficients are generally so much greater in the UV than in the IR that a UV method is to be preferred. To maximize the temperature rise, one could illuminate the wall with a narrow beam of radiation which slowly scanned the wall. The temperature rise can be sensed by a thermal-infrared detector that scans the wall in synchronism with the illuminating beam. The illuminating beam can be either chopped or unmodulated. Because the observed temperature change may be strongly dependent on which beam type is chosen, both methods will be examined.

b. Chopped-Radiation Methods

By means of a suitable chopper, the incident radiation intensity, I, as a function of time, t. can be made to have the form shown below:



Chopped Incident Radiation as a Function of Time

This function can be represented by the formula

$$I = \frac{1}{3} I_0 (1 + \cos \omega t) \tag{1}$$

consisting of a steady component

$$I_{g} = \frac{1}{2} I_{O} \tag{2}$$

and an alternating component

C

2

$$I_{g} = \frac{1}{2} I_{g} e^{i\omega t}$$
 (3)

where for mathematical convenience we have replaced $\cos \omega t$ by $e^{i\omega t}$.

If a lock-in amplifier tuned to the angular frequency ω is used with the detector, only the effect of the component I_a is observed. It is to be noted that since I_a takes on both positive and negative values, this component of the intensity implies formally that power is both put into and taken out of the wall.

If the thin absorbing layer is of thickness d and absorption coefficient α , at the wavelength of the incident beam, the intensity transmitted through the layer is

$$I_t = (1-R) I_a e^{-\alpha d} = \frac{(1-R)}{2} I_o e^{-\alpha d} e^{i\omega t}$$
 (4)

where R is the reflectance of the film at the irradiation wavelength and the power per unit area absorbed in the layer is therefore

$$W = \frac{(1-R)}{2} I_0 (1 - e^{-\alpha d})$$
 (5)

where the factor $e^{i\omega t}$ has been suppressed.

We assume that the transmitted intensity, given by Equation (4), is slowly absorbed by the underlying wall and therefore does not appreciably affect the temperature distribution set up in the wall.

The temperature distribution in the wall is given by the heat flow equation

$$K \frac{\partial^2 T}{\partial x^2} - C \rho \frac{\partial T}{\partial t} - i\omega C \rho T \tag{6}$$

where x is distance into the wall, and K, C, ρ are, respectively, the thermal conductivity, specific heat and density of the wall material.

The solution of Equation 6 is

$$T = T_o e^{-\sqrt{\frac{1\omega}{k}}} \times$$
 (7)

where k is the thermal diffusivity, given by

$$k = \frac{K}{C \cdot c} \tag{8}$$

and T_0 is the surface-temperature amplitude.

The heat flux density at x = p is

$$q = -K \left(\frac{\partial T}{\partial x}\right) = K \sqrt{\frac{i\omega}{k}} T_0$$

$$x = 0$$
(9)

This flux density must equal the power W given by Equation 5. Therefore, the temperature amplitude at the surface is

$$T_{o} = \frac{(1-R) I_{o} (1 - e^{-\alpha d})}{2K \sqrt{\frac{1\omega}{k}}} = \frac{(1-R) I_{o} (1 - e^{-\alpha d})}{2 \sqrt{\omega K C}}$$
(10)

c. Unmodulated-Radiation Method

In the case of an unchopped incident beam the surface temperature is controlled by the thermal spreading resistance r_1 into the concrete wall. This quantity is given by

$$r_1 = \frac{1}{\|I\| + K} \text{ (deg/watt)} \tag{11}$$

where a is the radius of the incident beam and K, as before, is the thermal conductivity of the concrete wall.

In addition there is another resistance due to thermal radiation from the heated surface. This effect can be represented by a thermal resistance \mathbf{r}_2 given by

$$r_2 = \frac{1}{4 \epsilon \sigma T^3 \Pi a^2}$$
 (12)

where ϵ is the thermal emittance of the wall and σ is the Stefan Boltzmann constant.

Let P be the power delivered to the absorbing layer by the incident beam. Then the temperature rise T is given by "Ohm's Law" for heat

$$T = (1-R) \quad rP = \frac{(1-R) (1 - e^{-\alpha d}) I_{o}}{\frac{K}{a} + 4 \epsilon \sigma T^{3}}$$
 (13)

where
$$\frac{1}{r} = \frac{1}{r_1} + \frac{1}{r_2}$$
 (14)

The sample calculations below show that the unmodulated-beam method is greatly be to preferred. Equation (13) applies to the case of a non-scanning beam. The temperature rise will decrease with increasing scan rate. In actual practice, a number of implementations of the precise methodology are possible. One would be to irradiate a relatively large patch of the wall and view it at the same time. The entire area could be surveyed by moving the irradiated patch and thermal viewer together. Another possibility would be to irradiate only a small moving spot (more radiation per unit area) and view a large patch over the time period necessary to irradiate the whole patch. A third possibility is to view only the irradiated small spot and move the viewer with the subject spot. The precise implementation of the scheme will depend on a number of engineering tradeoffs concerning the expected signal strength, time available to survey the area in question, and so forth.

Sample Calculations. Based on the formulas derived above, the AC and steady-state temperature changes were calculated using ultraviolet irradiation and thermal viewing on concrete and wood. The parameters of the calculation are given together with their sources in Table III-4. absorption coefficient used in this calcuation was for PETN at a wavelength near 2,000 angstroms. This wavelength is shorter than the UV irradiation source peak used for calculation and some effort would have to be made to match the two. The PETN data were obtained from the Journal of Physical Chemistry, 77 910, (1973) showing a value of the absorption coefficient of 5.6 x 10^4 cm⁻¹ between approximately 45,000 and 55,000 cm⁻¹ (near 2000A). TNT in the UV region (near 2325A) has an absorption coefficient similar to that of PETN. Most of the other parameters are somewhat variable depending on the source, as the materials (concrete and wood) are variable in composition. A spectrum of TNT was run in the infrared which shows a maximum absorption coefficient of about 7,000 cm⁻¹ between 6.2 and 6.7 µm. These data, which are expected to be typical of most of the compounds in the two regions, are the reason for our choice of the ultraviolet region for surface irradiation.

TABLE III-4. PARAMETERS USED IN SAMPLE CALCULATIONS

Parameter	<u>Value</u>	Source and Comments
Ruv	0.1	Estimate from the Infrared Handbook
α PETN	5.6 x 10 ⁴ cm ⁻¹	J. Phys Chem, <u>77</u> , 910 (1973)
α _{TNT} (peak)	$1.38 \times 10^5 \text{ cm}^{-1}$	ARLCD-TR-78025 (1978)
d	$0.56 \times 10^{-7} \text{ cm}$	Taken from 1 μ g/cm ² , ρ PETN = 1.77
Io	$5 \times 10^{-3} \text{ watts/cm}^{2*}$	Estimate by P. von Thuna, ADL
K _{concrete}	8×10^{-3} watts/cm deg C	AIP Handbook
Kwood	2×10^{-3} watts/cm deg C	AIP Handbook
Cconcrete	0.65 j/gm deg C	Marks Standard Handbook for Mechanical Engineers
Cwood	1.75 j/gm deg C	Handbook of Chem & Physics
^P concrete	1.6 g/cm ³	AIP Handbook
Pood	0.6 g/cm ³	API Handbook
a	1 cm	
ω	6.28 Hz	For 1 Hz chopping frequency
€ IR	0.9	Estimate from related materials
T	300°K	

Ľ

 $[\]sigma$, the Stefan Boltzmann Constant = 5.67 x 10^{-12} $\left(\frac{\text{watts}}{\text{cm}^2\text{deg}^4}\right)$

This value will depend on the precise wavelength, bandwidth, and geometry of UV optics.

Using formula (10) for modulated UV irradiation (PETN)

$$T_{o} = \frac{(1-R) (1 - e^{-\alpha d}) I_{o}}{2 \sqrt{KCP\omega}} = 0.006^{\circ}C \text{ (concrete)}$$

= 0.01°C (wood)

Using formula (13) for unmodulated UV irradiation (PETN)

$$\Delta T = \frac{(1-R) (1 - e^{-\alpha d}) I_o}{\frac{K}{a} + 4\epsilon\sigma T^3} = 0.3^{\circ}C \text{ (concrete)}$$

= 1.1°C (wood)

The values for TNT would be approximately the same as those for PETN as the listed value is at the peak of the absorption band which extends out to near 2800A°. The fact that the latter value are well within the measurement capabilities of commercially available thermal imaging equipment forms the basis for our conclusion that the method is technically feasible.

3. Analyte UV Absorption Characteristics

UV absorption data for the analyte PETN were presented in the preceding discussion. Data on each of the other analytes of interest was also assembled and is in Table III-5.

it should be noted that most of the data in Table III-5 are for solutions of the analytes since these properties are customarily measured and reported in that manner. It is not clear that solution data are reliable indicators of the strength or location of UV absorption bands of the solid samples because these characteristics are often strongly influenced by the solvent. However, laboratory comparison of UV absorption spectra of the analyte 2,4-DNP in acetonitrile solution and in a KBr pellet and showed that the wavelengths corresponding to maximum absorption and the calculated absorption coefficients are quite similar. To the extent that this is also the case for the other analytes (and depending on the solvents), the data in Table III-5 may, in fact, indicate that relative locations and strengths of UV absorption bands. In that case, the data in Table III-5 suggest the following:

- 1. With the exception of NG, all analytes exhibit comparable absorption strengths (i.e., ε and α in Table III-5 are on the same order of magnitude for all analytes). Thus, the proposed approach should be equally applicable to all analytes;
- 2. The absorption bands of all analytes are sufficiently close to 2500 Angstroms that an illumination source capable of providing sufficient power output over some wavelength range centered on or about this wavelength may permit detection of all analytes (except NG). Commercially available mercury sources which emit the characteristic 2537 Angstrom line may be suitable for this purpose.

TABLE III-5. UV ABSORPTION DATA FOR ANALYTES IN SOLUTION

Compound	Solvent	<u>(v)</u> <u>v</u>	E(L/mole cm)	$\alpha(cm^{-1})$	Reference
TNT	Ethanol II ₂ 0	2270 2320	19,500	1.38×105	1 9
2,4 Dinitrotoluene	C ₅ H ₁₂ Ethano1	2320 2405	15,850	1.32x10 ⁵	-
2,6 Dinitrotoluene	651114	2270	12,000	8.48×104	•
KDX	Nethanol Solid H ₂ O Ethanol	2360 2360 2400 2130	10,000 11,000 10,000 11,000	9.02×10 ⁴	
NHA	Acetonitrile	2000 2220 (not peak)	10,000 ~2,000	5.6 ×10" vi.1 ×10"	· «c
Nitroglycerine	1120	2760	10	70	7
Jetryl	Ethanol	2250	25,100	1.37×10 ⁵	٦
Dipheny lamine	Methanol or Ethanol Ethanol, 77°K	2850 5500	20,400 31,000	1.4 ×10 ⁵ 2.12×10 ⁵	- 5
1,3,5 Trinitrobenzene	50-50, Methanol-H ₂ 0 G ₇ H ₁₆ C ₇ H ₁₆ G ₇ H ₁₆ Ethanol	2250 2800 3500 2220 2250	25,700 550 182 31,600 25,700	2 ×10 ⁵	۷
2,4 Dinitrophenol	HoO Ethanol Ethanol	3600 2920 2530	17,800 9,100 10,200	1.63×105	· 21
s 7 molar absorptivity; α 3 · c	3 · : : : : : : : : : : : : : : : : : :				•

. A motar absorptivity; α E re

Table II-5 (Continued REFERENCES

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UV Absorption data for solutions of analytes were presented in Table III-6. We also obtained solid state UV absorption coefficients of six analytes to determine whether the tentative conclusions based on examination of solution data could be confirmed. A technique used extensively in infrared analysis—the preparation of KBr pellets containing the analyte—was attempted and found to be satisfactory through the Cary 219, 400-200nm, ultraviolet range. Macropellets of 13-mm diameter were pressed under 20,000 pounds pressure, using oven-dried KBr to prevent moisture from clouding the pellet.

Initial attempts to use a Wig-L-Bug device to thoroughly mix sample and KBr proved unsatisfactory—inexplicable clouding occurred, even in thoroughly dried containers. A successful method finally used involved gentle but thorough grinding in a mortar and pestle. Some experimentation was necessary to determine the right concentration to yield a measurable peak.

Two methods of measurement were used, both giving comparable results. The first method involved plotting an air vs. air baseline, then obtaining both blank KBr pellet and sample KBr pellet traces. At the sample peak, the blank reading was subtracted. In the second method, a baseline was obtained with blank KBr pellets in each beam. When the sample pellet was substituted on the sample side, the absorbance at the peak could be read directly.

Thickness of the pellets was measured with a micrometer and calculations made as noted in Table III-6. These findings confirm the tentative conclusions noted earlier.

4. Laboratory Demonstration.

The thermal imaging equipment required for laboratory demonstration of the practical utility of the UV illumination/thermal imaging sampling protocol was rented from Inframetrics, Inc. for one week. Using this equipment and UV illumination sources already on hand in Arthur D. Little, Inc. laboratories, we were able to observe the presence of small quantities of solid analyte on all of the available surface types of interest except brick. However, the observed contrast between analyte and surrounding surfaces was less than would be desirable for routine field use. Independent measurements indicated that the power output of the UV illumination sources used was on the order of 150 microwatts to 1 milliwatt, which is less than that assumed in calculations above. Substitutions of the experimental values for UV power output in those calculations results in a predicted temperature rise on the order of that which was, in fact, observed. Thus, while these experiments confirm the practical utility of the method, they also indicate that for optimum performance a UV illumination source having a higher power output in the desired wavelength region should be used. We expect that a UV source with the desired performance characteristics and operational characteristics that would permit its safe use under field conditions can be obtained commercially.

TABLE III-6. SOLID STATE ANALYTE UV ABSORPTION COEFFICIENTS

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	TABLE	TABLE III-6, SOLID SIAIE ANALIIE OV ABSONIIION COLITICIENTO	D SIAIE ANAL	ILE UV ABSUN	TION COPIL			
Sample	Wt. Samplo mg	Density g/cc_	Wc. KBr mg	Density 8/cc	л Мах	Abs.*	Thickness	$\alpha(cm^{-1})$
1,3,5-TNB		1.688	324.888	2.75	230	$\frac{1.65^{(2)}}{1.70^{(1)}}$.935	1.21×10^5 1.25×10^5
2,4-DNP	650.	1.683 ²⁴	199.74	2.75	262	2.62(1)	09.	.90 × 10 ⁵
2,4-DNT	050.	1.521	320.111	2.75	264	1.52(1)	.91	.59 × 10 ⁵
RDX	. 446	1.82 ²⁰	319.654	2.75	240	.86 ⁽¹⁾	06.	.045 x 10 ⁵
2,4,6-TNT	т .136	1.654	260.114	2.75	240	. 54 (2)	.73	.085 × 10 ⁵
Tetryl	.127	1.5719	250.179	2.75	228	. 38(2)	69.	.062 × 10 ⁵

*Correction made for KNr absorbance by subtraction or using dual beam method with blank KBr in reference side.

- (1) Subtraction method.
- (2) Dual beam merhod.

E. ANALYTE DETECTION USING UV IRRADIATION AND UV PHOTOGRAPHY:

1. Background.

UV light directed at analytes present on a surface may be 1) transmitted, 2) reflected, or 3) absorbed. Measurement of the temperature rise resulting from UV absorption is the principle underlying the thermal imaging approach. Any UV absorption will necessarily be accompanied by a decrease in the UV reflectance, and observation of the latter quantity may also be a feasible detection method. In this case, surfaces on which analytes are present would appear as dark areas against a lighter background when illuminated with a UV source and viewed with a UV selective detector. The detector could consist simply of black and white film in a large format camera equipped with a UV filter centered on or about 2500 Angstroms. The advantages of this approach would include the simplicity and ease of operation of the required equipment and the fact that the resulting photographs could be maintained as a permanent record of the investigations.

Preliminary Experiments.

Two types of experiments were performed to assess the practical utility of this approach. In one experiment, the diffuse UV reflectance from a concrete surface to which 60 µg/cm² of 2,4-DNP were added was measured in a UV spectrophotometer. About a 59% reduction in reflectance at 3600 Angstroms was observed. This observation suggested that for concrete and probably for other surface types as well, the anticipated effects were, in fact, observed and that the measurements could probably also be made at much lower analyte concentrations.

In a second set of experiments, UV photographs of several analytes spiked on surfaces were obtained. Figure III-7 is a photograph showing the contrast obtained for 2,4-dinitrophenol, tetryl, 2,6-dinitrotoluene, RDX and TNT in 1.3 x 10^{-4} g/cm² amounts on a thin layer chromatography silica gel substrate. The photograph was obtained using a 2400 A filter and a Ziess HBO 200 W source with the glass lens removed.

Figure III-8 shows the concentration sensitivity for differing amounts TNT, tetryl and 2,4-dinitrophenol on concrete using a 3650 A filter. The top row has concentrations of approximately 6×10^{-5} g/cm² for each material (including the amount lost by diffusion into the substrate). The second row has about twice the amount of TNT and half the amounts of tetryl and 2,4-dinitrophenol although the spot size is somewhat variable.

Characteristics of the UV irradiation and viewing filter would have to be optimized for maximum contrast. Additional experiments using different UV illumination sources and narrow bands pass UV filters were performed to establish the optimum conditions for UV photography.

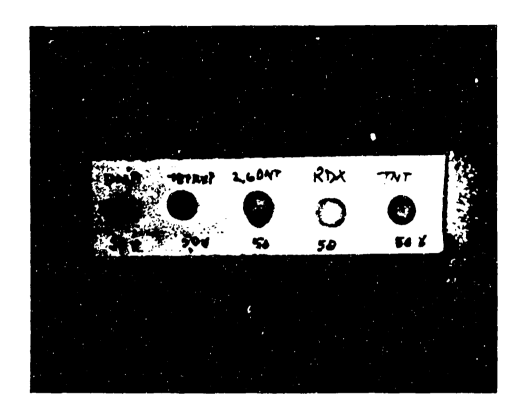


Figure III-7. UV Photography of 130 mg/cm² Each of 2,4-DNT, Tetryl, 2,6-DNT, RDX, and TNT on TLC Plate

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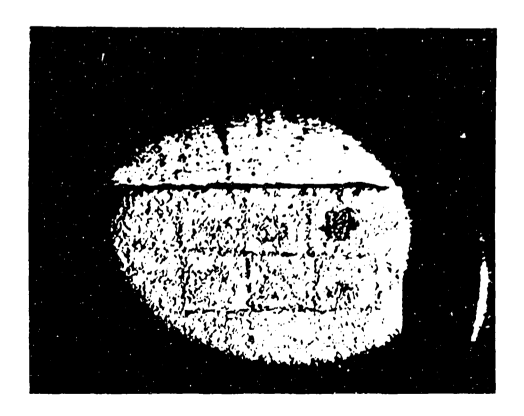


Figure III-8. UV Photography of TNT, Tetryl, and 2,4-DNP on Concrete

The camera and lens for these experiments consisted of long bellows Bausch and Lomb microscope camera that could be converted to a horizontal position. The camera has a variable shutter to which we could attach a simple, 250 mm focal length, quartz lens. A variable iris was used to reduce the aperture and thus effectively increase the depth of field. This procedure compensated for the change in effective focal length encountered when focusing with visible light then photographing at a shorter wavelength, thus throwing the image out of focus.

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This camera has a 4 x 5 sheet film format with which we used Polaroid P/N 55 sheet film providing both the positive print and a negative. The film is a slow high resolution type film with apparent adequate sensitivity to the ultraviolet to allow a relatively short exposure time. Five seconds was adequate with the Leiss lamp.

The experiments described previously utilized a high preasure mercury arc (Zeisch HBO 200 Watt) which produces a substantial amount of energy centered on the 365 nm mercury emission line. This radiation was selectively transmitted to the camera lens with the use of a barrier filter with a maximum transmission at 365 nm.

In additional experiments, a Mineralite Model S-61 source was obtained for evaluation. This is a low pressure mercury lamp which is reported to have a higher intensity of 254 nm radiation. This lamp is of particular interest as a radiation source for the thermal imaging experiments where it is essential to optimize the absorption of radiation to achieve the maximum subsequent thermal emission. With this lamp as a light source, a series of photographs were taken of the test spots of explosives on various substrates using different barrier filters. Experimentally it was determined that a 10-minute exposure was required to produce an adequate image using a 254 nm hand pass barrier filter over the camerallens.

The results of these experiments are shown in Table III-7.

These data indicate that it takes a much longer exposire interval to obtain a suitable image with the shorter 254 nm radiation. This would not be a significant handicap if there were a distinct advantage to be gained. However, we observe approximately the same level of gray image for each of the representative substrates for both radiation sources. In order to successfully photograph any explosive residue, it is obvious that the image must appear darker than the background or substrate material. In our evaluation we have applied the explosive compound to silica thin layer chromatography substrates as a standard of comparison. A good contrast image was obtained for most of the explosives on this substrate and 50 microgram spots could be detected for most of the compounds. Nitroslycerine and PETN did not absorb at this wavelength and do not phocograph well under these conditions.

Ultraviolet Products, Inc., South Pasadena, CA

TABLE III-7. RESULTS OF UV PHOTOGRAPHIC EXPERIENENTS

Radiation Source	High Pressure Hg Arc Hg Arc, Zeiss	Low Pressure Hg Arc, Mineralite
Most intense UV radiation	365 nM	254 nM
Optimum barrier filter for camera	365 nM	254 nM
Explosure level (film response)	5 seconds	10 minutes
Substrate Response:		
Silica, Chromatograph plate	White	White
Concrete block	Light gray	Light gray
Transice	Medium gray	Medium gray
Galvanized stell	Reflective	Reflective
Brick	Dark gray	Dark gray
Wood	Dark gray	Dark gray
Glass		Black

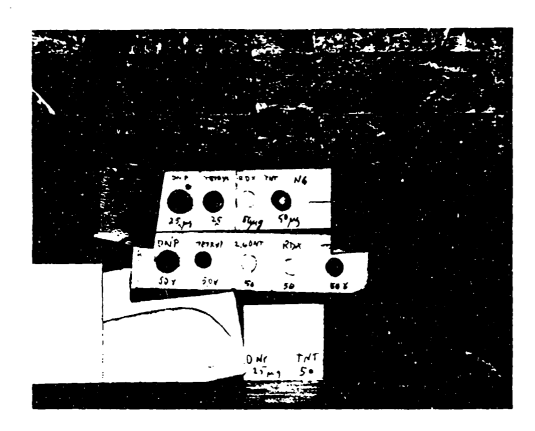
Typical examples of these results are shown in attached photographs. These photographs were taken using the low pressure mercury source with an exposure of 10 minutes. Figure III-9 was taken with the 365 nm transmission filter over the lens. We note excellent contrast for DNP, tetryl and TNT. Figure III-10 shows the effect of using the 254 nm tansmission filter which changes the focal length with the shorter wavelength. It is interesting to note that the DNP and tetryl have less contrast at this shorter wavelength, while RDX has slightly better contrast.

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Figure III-10 includes a sample of transite containing 50 micrograms quantities of applied explosives. Recognizable spots are noted for DNP, tetryl, RDX and TNT while TNB is barely visible.

It has been reported that normal optical glass will transmit enough 365 nm radiation that suitable photographs can be produced with a conventional camera lens. However, this would be a less than optimum condition and totally ineffective for the 254 nm radiation and therefore not investigated.

Taken together, these results suggest that further investigation of this approach appears to be warranted. In particular, it would be desirable to evaluate under actual field conditions the ability of this approach to detect those analytes for which these experiments indicate it is best suited.



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Figure III-9. Analytes on Silica Gel and Metal. 365 nm

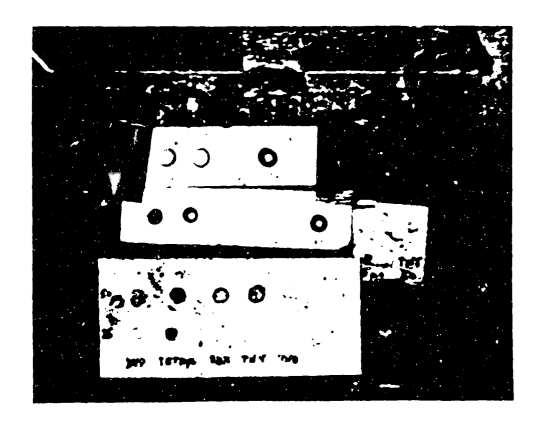


Figure III-10. Analytes on Silica Gel (top) and Transite (bottom). 254 nm

IV. QUANTITATIVE METHODS DEVELOPMENT

A. INTRODUCTION

The quantitative method selected for developmental testing involved solvent extraction using alternative procedures to conventional wipe or swab methods. Existing USATHAMA methods for the determination of explosives, modified as necessary for this particular application, were used for analysis of the resulting extract.

The objective of this testing was development of procedures for quantitative determination of specific compounds down to concentrations as low as 5 $\mu g/10~cm^2$. The approach used to achieve this objective involved in each case the spiking with known amounts of analytes of new, uncontaminated samples of each of the surface types of interest obtained from building materials dealers. The samples of conductive non-sparking flooring that were available for this study were known to have been contaminated with explosives/explosive residues, thereby precluding the spike and recovery approach. The results obtained by applying the procedures developed for other surface types to samples of this material are described in Section IV.C.3.

At the direction of the Technical Project Officer, emphasis was placed on the development of procedures for organic analytes. Methods based on an extraction approach similar to that described in this report for organic compounds are used widely for the determination of inorganic species, including those of interest in this study, in soils, sediments, sludges, etc. No methods for inorganic species which would represent substantive improvement over those methods were finally identified.

B. SEMIQUANTITATIVE CERTIFICATION TESTING

Preliminary semiquantitative certification was accomplished by spking analytes into the quantity of acetonitrile specified in the existing USATHAMA method prior to any concentration or solvent exchange steps, and continuing through the steps of the analytical method. The certification test involved analyzing five analyte concentrations plus a blank, one time each on three or more days. The resulting data is summarized in Volume II of this report.

Tables IV-1 and IV-2 summarize the analytical method conditions and the statistical data, respectively. (Tables IV-1 and IV-2 are reproduced as Tables I-1 and I-2 in Volume II of this report).

C. QUANTITATIVE CERTIFICATION TESTING

1. Introduction

Quantitative certification was accomplished as noted above by spiking analytes in acetonitrile solution directly onto new, uncontaminated samples of each of the surface types of interest, extracting the spiked

<u>.</u>	Flou Rate	l mi./min	1 mf./min	1 ml./mln	30 mt./min	30 mt/min	30 ml./min	30 ml./mtn	1 ml./min	1 m./m1:	1.0 at./atn	1.0 m./min	1.0 ml/min	1.0 ml/mtn	1.0 ml./min	1.0 ml/min	1.0 ml./min
	Sulvent System/ Carrier Gae	65/35 CH JCH/H ₂ 0	35/65 CH ₃ CH/ ,005H t-butyl	ammontum hydroxide	5% Methane/argon	5% Methane/argon	5% Hethane/argon	5% Metliane/orgon	90/10 methanol/H ₂ 0	60/40 methano1/11 ₂ 0	0.08M neets asid adjusted to pll 3.1 with ammonium hydroxide/Cll gCN	0.08M acetic acid adjusted to pll 3.1 with ammonium hydroxide/Cil ₃ CN	0.08M acetic acid adjusted to pll 3.1 with ammonium hydroxide/Cil ₃ CM	0.08M acetic acid adjusted to pH 3.1 with ammonium hydroxide/CH ₃ CM	0.08M acetic acid adjusted to pli 3.1 with ammonium hydroxide/Cii ₃ CN	0.08H acetic acid adjusted to pll 3.1 with ammonium hydroxide/(3) ₃ CN	0.08N acetic acid adjusted to pH 3.1 with assumium hydroxide/CH ₃ CN
	Retrution Time	6.2 mln	24.6 mtn	30.5 min	19.90 min	16.20 min	30.95 mtn	28.14 min	230 вес	416 sec	12.5 ∎In	13.5 mIn	17.2 min	24. 3 a fn	25.6 min	26. i safın	39.2 min
ri carf the B NDITIONS	Property and a second	Temp or Solvent Isocratic	faceratic	Isocratic	100C for 6 min 15C/min to 165C:	hold 8 min 15C/min to 200 Cı	hold 6 min	£	laceratic	factatic	Inttial: 30% CU ₃ CM Final: 50% CU ₃ CM Time: 35 min Gradient: linear	Prittal: 30% CH ₂ CN Final: 50% CH ₂ CN Time: 35 min Gradient: linear	Initial: 30% CH CN Final: 50% CH CH Time: 35 min Gradient: 11near	Initial: 30% CH ₃ CN Final: 50% CH ₃ CN Time: 35 min Gradient: 11near	Initial: 30% CN ₃ CN Final: 50% CN ₃ CN Time: 35 min Grodlent: Finear	Initial: 30% Ch ₁ Ch Final: 50% Ch ₂ Ch Time: 35 min Gradlent: linear	Intefal: 30% CH CH Final: 50% CH CH Time: 35 min Gradient: Tinear
ANALYTICAL METHOD CONDITIONS		Pellicular LC-18, 40n, 50 x 4.6 am	Pellicular LC-18, 40μ, 50 x 4.6 mm	Pelifeular 1.C-18 40µ 50 x 4.6 mm	None	None	None	None	None	None .	Pellfular I.C-19,40µ 50 x 4.6 mm	Pelifeular I.C-18, 40µ 50 x 4,6 mm	Pellicular LC-18,40µ 50 × 4.6 mm	Pellicular 1.6-18, 40µ 50 x 4.6 mm	Pellicular LC-18, 40µ 50 x 4.6 pm	Pelifcular I.C-18, 40µ 50 x 4.6 mm	Pellicular 1.C-18, 40µ 50 × 4.6 mm
A. A. A.	en l'o	Spherisort ONS, SH, 250 x 4.6 mm	Splierinorb (4)5, 511, 250 x 4.6 mm	Spherinoxb 00S, 5µ, 250 x 4.6 mm	3% 0V-225 on 100/120 Cas Chrostly x" x 2 mm 1D x 6' glass column	3% 09-225 on 100/120 Cae ChromQ ½" x 2 mm 1D x 6° glass column	3% OV-225 on 100/120 Gas ChromQ %" x 2 cm 1D x 6' glass column	32 0V-225 on 100/120 Gas ChromQ \(\frac{1}{4}\) x 2 mm ID x 6" glass column	U Bondpack C18 fam x 30 cm	U Roudpack C18 4mm x 30 cm	Sphertsorb (1915, 511, 250 x 4.6 me	Sphertsorb 005, 5p, 250 x 4.6	Sphertsort Obs, 5µ, 250 x 4.6 mm	Sphertsorb 005, Su. 250 x 4.6	Spliertsorb ODS, 5µ, 250 x 4.6 mm	Sphertsorb 008, 5p, 250 x 4.6 mm	Spherfsorb OBS, 5p, 250 x 4.6 mm
!		IIP1.C	HP!,C	HPLC	၁၁	99	25	8	HPLC	IIPLC	url.c	HH.C	mr.c	EPLC.	9. 1. 1.	3710	BPLC.
	1	PETN	2, 6-11HT	- ÿ	/2,4-hNT	2,6-DNT	1, 3, 5-THB	Z,6,6-1NT	1).V	Tetryl	·iva	RDX	T N T	2,4-INT	TNT	Tetryl	N.Y

TABLE IV-2. SEMIQUANTITATIVE CERTIFICATION TESTING STATISTICAL DATA SUMMARY

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	Int. Reference	0,450 9	9.888	2119.5 B	0.634 5	-0,012 5	0.007 5	-0.009 5	-0.634 3	-0.004 5	5.397 10	0, 698	0.674 10	1.585 10	0.269 10	-5,410 10
	Stope It	1.046 0.	1,061	0,939 21	0.743 0.	0- 690.1	0.058	1.054 -0,	0- 386 -0.	0.862 -0.	0.971 5.	1,009	0 366.0	0.990	1,012 0	1.0505
Corr.	Caril.	0.996	946	666.0	0.967	0.989	666.0	676.0	6.69	6.995	966.0	0, 398	666.0	0.999	666.0	666'0
Det.	Libit	1.77 ug/ml.	0.26 ug/el.	. 4.54 ug/ml.	0.11 ug/ml.	0.09 ug/ml.	0.12 ug/al.	0.12 ug/mt	12.25 ug/ml.	0.08 ug/ml.	30 ng/mL	70 ng/ml.	25 ng/ml.	24 ng/mL	24 ng/mi.	25 ng/ml.
USATILAM	Hethod	none	HRI Hethod	MR1 Method	246 TMT-WA-02	246 THT-UA-02	246 TNT-MA-02	246 TNT-44-02	DPA-WA-01	TETRYL-WA-02	HRI Nethod	MIR Nethod	MKI Method	MRI Method	PRI Method	MKI Method
4	Volume	70 uL	70 ul.	70 ul.	, ut.	Jo C.	ן ה	- -	200 ut	200 ut	100 ul.	100 nL	100 al.	100 of	100 ul.	100 ml.
Chart	Speed	0.1 In/mIn	0.1 tu/atn	0.1 In/mIn	0.5 cm/min	0.5 cm/min	0.5 cm/msn	0.5 cm/min	0.5 cm/min	0.5 cm/min	0.1 in/min	0.1 In/#\$a	0.1 In/min	0.1 In/min	0.1 In/min	0.1 tu/min
	Atto.	0.01 AUPS	O.OL AUPS	0.01 AUFS	1×10-11×8	1x10-11x8	1x10-11x8	1×10-11x8	0.1 AUFS	O.1 AUFS	0.01 AUFS	O.O. AUP'S	0.01 ABI'S	0.01 AUFS	0.01 AGES	0.03 2005
	Detector	UV at 230 am	UV at 230 mm	UV at 230 nm	ECD #30MC	ECD 63000	ECD #300C	ECD @30HC	UV at 254 nm	UV .AC 254 AM	UV ot 254 nm	(IV at 254 nm	UV at 254 nm	11V LE 254 MB	UV at 254 nm	UV .1. 254 118
	Analyte	FESTS	7.6-UNT		, , 4-INT	. 6-DHJ	1.1.5-THB	1.4.6-TWF	V.10	letryl	ME	KDX	T'NB	2,4-DNT	TWI	Terryl

samples with acetonitrile using ultrasonic agitation, and analytical procedures used for preliminary certification. The analytical methods that we developed were evaluated according to the procedures specified in the 1980 USATHAMA QA Plan. The resulting data is summarized in Volume II of this report.

2. Analytical Methods

The analytical procedures that were developed and evaluated are described in detail on pages 76 through 87 of Volume I of this report. (These analytical methods are also presented on pages 46 through 57 of Volume II of this report.

3. Results and Discussion

Table IV-3 summarizes the statistical data obtained from Quantitative Certification Testing. (Table IV-3 is reproduced as Table 1I-1 in Volume II.)

As noted above (Section IV.A), the samples of <u>conductive non-sparking</u> flooring that were available for this study were known to have been contaminated with explosives/explosive residues, thereby precluding quantitative certification testing. Instead, the analytes of interest were determined in these samples by the methods described in Section III.C.2.

Two samples of conductive non-sparking flooring from an operating explosives building were received from Dr. Harold J. Matsuguma, Chief, Chemistry Branch, Energetic Materials Division, ARADCOM. One sample consisted of a single large sheet about two square feet in area which reportedly represents a commonly used roll material. The other sample consisted of several small chunks which reportedly represent another conductive coating which is trowelled onto a floor and allowed to harden.

Of the analytes of interest in this study, these samples were reportedly exposed to RDX, TNB, TNT, tetryl, and PETN. Two subsamples, each approximately $10~\rm cm^2$ in area, of each of the two flooring types were extracted with acetonitrile using ultrasonic agitation and the analytes RDX, TNB, TNT, and tetryl were determined in the resulting extracts. The results of these analyses are presented in Table IV-4.

Analysis of wood samples spiked with explosives using these analytical methods met with mixed results. For many samples, analyte detection limits and recoveries comparable to those obtained for other analyte-surface combinations were obtained. However, in certain samples large interfering peaks which precluded identification and quantification of the analytes of interest were observed. These interferences appeared unpredictably among subsamples taken from a single piece of kiln-dried dimension lumber obtained from a local building materials dealer, and are presumed to be due to naturally-occurring compounds distributed

TABLE IV-3. QUANTITATIVE CERTIFICATION TESTING STATISTICAL DATA SUMMARY

	Metal			rete	Bri	ck	Transite		
	D.L. ug/cm ²	% Rec	D.L. ug/cm ²	% Rec	D.L. ug/cm ²	% Rec	D.L. ug/cm ²	% Rec	
DNP	0.33	96%	1.74 (0.35	31% 34%) ²	1.59	55%	2.22 (1.15	34% 30%) ²	
RDX	0.25	96%	0.63	78%	2.11	72%	3.48 (0.80	67 % 82%) ²	
TNB	0.28	95%	0.98	75%	2.12	74%	3.46 (0.76	59% 71%) ²	
2,4-DNT	0.90	84%	1.08	78%	2.10	69%	3.52 (0.62	65% 79%) ²	
2,4,6-TNT	0.60	97%	1.57	74%	1.68	66%	3.18 (0.68	51% 60%) ²	
Tetryl	1.95	88%	4.29	51%	2.60	68%	4.13	56%	
DPA	0.50	94%	2.44 (1.15	74% 84%}	2.14	69%	3.71 (0.90	67% 82%) ²	
2,6-DNT	2.85	94%	6.46	86%	6.36	49%	2.04	79%	
NG	9.36	94%	21.7	76%	32.5	44%	26.1	72%	
PETN	10.2	86%	5.39	83%	20.6	62%	10.0	82%	

Calculated from four days of target vs. found concentrations using the proceduces specified in the 1980 USATHAMA QA Plan.

² Calculated from three days of target vs. found concentrations.

QUANTITATIVE METHOD FOR THE DETERMINATION OF DNP, RDX, TNB, 2,4-DNT, TNT, TETRYL, AND DPA ON SURFACES

1. Application

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Method used to extract the following compounds from metal, brick, concrete, transite surfaces:

2,-,-dinitrophenol DNP
cyclotrimethylenetrinitramine RDX
1,3,5-trinitrobenzene TNB
2,4-dinitrotoluene 2,4-DNT
2,4,6-trinitrotoluene 2,4,6-TNT
2,4,6-trinitrophenylmethylnitramine Tetryl
diphenylaminu DPA

A. Tested Concentration Range

DNP	0.25	ug/cm ²	_	5.00	່ນg/cm²
RDX	0.25	ug/cm'	_	5.00	ug/cm:
INB		ug/cm ²			
2,4-DNT	0.25	ug/cm ²	_	5.00	ug/cm²
THT	0.25	ug/cm2	-	5.00	աg/cm²
Tetryl	0.25	μg/cm ²	-	5.00	µg/cm ²
DPA	0.25	ug/cm²	-	5.00	ug/cm2

B. SerialLivity

Instrument response for each analyte is given below:

Analyta	Concentration	Response			
DHP	25.04 ng/mL	3700 area unita			
KDX	24.86 ng/ml.	1388 area units			
7 413	24,97 ng/mL	4437 area unite			
2.4 -DNI	24,99 ng/ml.	6090 area unita			
1101	24.91 ng/ml.	4104 ares units			
1 cryl	25.30 ng/mL	2895 area unic			
DFA	24.91 ng/mL	2101 area unite			

G. Detection Limit

Bue lable 11-1.

D. Interfyjon ar

I effective present in some bifix and transite samples were apparently under tather than systematic. For example, the HPLC smallysis of extraction B of one blank (unspiked) brick surface (extracted for 9)

indicated the presence of a compound with a retention time of 1510 seconds. This compound interfered with the TNT. However, this interference was not observed in any other blank brick sample. Therefore, only the brick samples analyzed on Feb 9 were corrected for the interference.

E. Analysis Rate

Six samples can be extracted and prepared for analysis in three hours. Rate of analysis is given below, excluding calibration standards:

DNP	8	samples	1n	an	8	hour	day
RDX	8	samples	in	an	8	hour	day
TNB	8	samples	in	an	8	hour	day
2,4-DNT	8	samples	in	an	8	hour	day
TNT	8	samples	1n	an	8	hour	day
Tetryl	8	samples	in	an	8	hour	day
DPA	8	samples	in	an	8	hour	day

2. Chemistry

2,4,-dinitrophenol C6H4N2O5 CAS RN 51-28-5 MP 112-114C

Cyclotrimethylenetrinitramine C3H6N6O6 CAS RN 121-82-4 MP 205-206C

1,3,5-trinitrobenzene C6H3N3O6 CAS RN 99-35-4 MP 122.5C

2,4-dinitrotoluene C7H6N2O4 CAS RN 121-14-2 MP 71C

2,4,6-trinitrotoluene C7H5N3O6 CAS RN 118-96-7 MP 80.1C

2,4,6-trin1troph@nylmethylnitramine CAS RN 479-45-8 MP 130C Explodes 187C

41phanylamine C17411N CA5 RN 122-39-4 MP 53-54C BP 307C

3. Apparatus

A. Instrumentation

Waters Associates Model 6000A Solvent Delivery System Waters Associates Model M-45 Solvent Delivery System Waters Associates Model 660 Solvent Programmer Waters Associates Model 440 Absorbance Detector Waters Associates Intelligent Sample Processor (WISP) Spectra-Physics Minigrator Hewlett Packard 7133A Recorder

B. Parameters

Column: Spherisorb ODS 5μ , $250 \times 4.6 \text{ mm}$ ID Precolumn: Pellicular LC-18, 40μ , $50 \times 4.6 \text{ mm}$ ID

Solvent System: linear gradient

Initial: 30/70 CH₃CN/0.08 M acetic acid adjusted

to pH 3.1 with NH40H

Final: 50/50 CH₃CN/0.08 M acetic acid adjusted

to pH 3.1 with NH4OH

Time: 35 minutes Detector: UV at 254 nm Flow Rate: 1.0 mL/min Attenuation: 0.01 AUFS Injection Volume: 100 μ L

C. Hardware/Glassware

Westinghouse Ultrasonic cleaner
8 ounce jars with teflon lined caps
25 mL graduated cylinders
microliter syringes
volumetric flasks - 50, 10, 5 mL
vials - WISP and 14 mL, with teflon lined caps

D. Chemicals

acetonitrile, HPLC grade nitrogen acetic acid ammonium hydroxide Standard Analytical Reference Material for each analyte

4. Standards

A. Calibration Standard

Stock Solution A: DNP, RDX, TNB, 2,4-DNT, TNT, TETRYL, DPA Prepare individual stock solutions of 5.0 mg/mL. Combine 500 μ L individual stocks and dilute to 5 mL.

Stock Solution B:

Dilute 625 μ L Stock Solution A to 5 mL. Concentration is 62.5 μ g/mL of each analyte

Calibration standards prepared in 25 mL volumetrics adding $\rm H_20$ so final solution is 55% $\rm H_20/45\%$ CH₃CN:

Cal Std.	uL Stock B Added	Concentration Each Analyte
1	10	25.0 µgL
2	20	50.0 μgL
3	40	100.0 ugL
4	80	200.0 ugL
5	200	500.0 ugL
6	400	1000.0 ugL

B. Control Spikes

Spiking stock solutions were prepared using 0.5 mg/mL stock solution.

ul Stock Soln/x mL CH3CN	Concentration Each Analyte
125 uL/10 mL	6.2 mgL
250 µL/10 mL	12.0 mgL
500 µL/10 mL	25.0 mgL
500 μL/5 mL	50.0 mgL
625 µL/5 mL	62.5 mgL
1250 µL/5 mL	125.0 mgL
2500 µL/5 mL	250.0 mgL

400 μ L 6.2, 12.5, 25.0, 62.5, 125.0 mgL Stocks spiked onto 10 cm^2 concrete and metal surfaces.

 $200~\mu L$ 12.5, 25.0, 50.0, 125.0, 250.0 mgL stocks spiked onto $10~cm^2$ brick and transite surfaces.

Concentration of analytes on surface after spiking: 2.5 μ g, 5.0 μ g, 10.0 μ g, 25.0 μ g, 50 μ g.

5. Procedure

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Extraction A

1. Spike 10 cm² surface sample with acetonitrile spike solution (volume dependent on surface type). Allow solvent to evaporate.

- 2. Transfer sample to 8 ounce jar and add 20 mL CH₃CN. Cover jar with teflon lined cap.
- 3. Sonicate for 10 minutes.
- 4. Transfer extract A to 50 mL volumetric flask, add 27 mL 0.08 M acetic acid and bring to volume with CH₃ CN. Save surface sample for extraction B.

Extract A ready for analysis.

Extraction B

- 1. Add 20 mL CH₃CN to jar with surface sample. Sonicate for 10 minutes.
- 2. Transfer surface sample to a second jar. Add 20 mL $\rm CH_3CN$ and sonicate for 10 minutes more.
- 3. Combine extracts from steps 1 and 2 and evaporate using nitrogen to less than 5 mL.
- 4. Transfer evaporated extral to 10 mL volumetric flask. Add 5.5 mL to 0.08 M acetic acid and bring to volume with CH₃CN.

Extract B ready for analysis.

6. Calculations

Calculate found concentration for each analyte in each sample extract from daily calibration data.

Multiply found concentration by extract volume to find total ug in extract. Combine total ug in extracts A and B to find total ug on surface.

7. References

Lakings, D.B., Baker, R.J., and Ctook, M.V.. "Precision and Accuracy Assessment of the High Performance Liquid Chromatographic Analytical Technique for the Determination of Dinitrophenol (ANP); Cyclotrimethylone trinitramine (RDX); 1,3-Dinitrobenzene (DNB); 1,3-5-Trinitrobenzene (TNb); 2,4-Dinitrotoluene (2,4-DNT); Trinitrotoluene (TNT); 2,4,6-Trinitrophenyl-mathylnitramine (Terryl); and Diphenylamine (DPA)", Midwest Research Institute Technical Veport No. 1, USATHAMA Contract No. 1AAPKA1-81-C-0007, March, 1981.

QUANTITATIVE METHON FOR THE DETERMINATION OF 2,6-DNT AND NG ON SURFACES

1. Application

Method used to extract the following compounds from metal, brick, concrete, transite surfaces:

2,6-dimitrotoluene 2,6-DNT nitroglycerine NG

A. Tested Concentration Runge

2.6-DNT 1.00 µg/cm² to 20.00 µg/cm² NG 12.50 µg/cm² to 125.00 µg/cm²

B. Sensitivity

Instrument response for each analyte is given below:

Analyte	Concentration	Response
2,6-0VT	0.10 pg/mL	208750 area units
NG	1.25 pg/mL	186010 area units

C. Detention Limit

See Table II-1.

D. Interfarences

No interferences were observed

E. Analysis Rate

Six samples can be extracted and prepared for analysis in three hours. Rate of analysis is given below excluding calibration standards:

2,5-DNT 16 Samples in an 8 hour day NG 16 samples in an 8 hour day

2. Chemistry

2,6 dinitrotoluere C7H6N2O4 CAS RN 606-20-2 MP 66C

MP Stable form 13.50

3. Apparatus

A. Instrumentation

Beckman Model 110A Solvent Metering Pump waters Associates Model 450 Variable Wavelength Detector Waters Associates Model U6K Injector Hewlett Packard 3390A Integrator/Recorder

B. Parameters

Column: Spherisorb ODS, 5μ , $250 \times 4.6 \text{ mm}$ ID

Precolumn: Pellicular LC-18, 40 μ , 50 x 4.6 mm ID

Solvent System: 35/65 CH₃CN/0.005 m t-butyl ammonium hydroxide,

pH 6.5. adjusted with 1N H3POu

Detector: UV at 230 nm Flow Rate: 1.0 mL/min Attenuation: 0.01 AUFS Injection Volume: 100 µL

C. Hardware/Glassware

Westinghouse Ultrasonic cleaner 8 ounce jars with teflon lined caps 25 mL graduated cylinders microliter syringes volumetric flasks - 50, 10, 5 mL vials - WISP and 14 mL, with teflon lined caps

D. Chemicals

Acetonitrile, HPLC grade nitrogen phosphoric acid t-butyl ammonium hydroxide

Standard Analytical Reference Material for each analyte.

4. Standards

A. Calibration Standards

Prepare individual stock solutions: 5 mg/mL 2,6-DNT 50 mg/mL NG Stock Solution A: Combine 200 μ L 2,6-DNT stock and 250 μ L NG stock and dilute to 10 mL CH₃CN. Concentration is 0.1 mg/mL 2,6-DNT and 1.25 mg/mL NG.

Calibration standards prepared in 10 mL volumetric flasks adding 50% H₂O/50% CH₃CN.

Cal Std.	μL Stock A added	2,6-DNT	NG		
1	10	0.1 ug/mL	1.2	μ g/mL	
2	20	$0.2 \mu g/mL$	2.5	µg/mL	
3	40	$0.4 \mu g/mL$	5.0	μg/mL	
4	80	0.8 µg/mL	10.0	ug/mL	
5	200	2.0 µg/mL	25.0	μg/mL	

B. Control Spikes

Spike Solutions prepared following chart below:

Spike Solution	Amount Stock	Dilute with CH ₃ CN to	Concentration
1	1 mL of 5 mg/mL 2,6-DNT	5 mL	1 mg/mL 2,6-DNT
2	0.5 mL of 5 mg/mL 2,6-DNT	5 mL	0.5 mg/mL 2,6-DNT
3	0.5 mL of 5 mg/mL 2,6-DNT 0.625 mL of 50 mg/mL NG	5 mL	0.5 mg/mL 2,6-DNT 6.25 mg/ml NG
4	0.250 mL of 5 mg/mL 2,6-DN7 0.312 mL of 50 mg/mL NG	5 mL	0.25 mg/mL 2.6-DNT 3.12 mg/mL NG
5	0.200 mL of 5 mg/mL 2,6-DN7 0.250 mL of 50 mg/mL NG	5 mL	0.20 mg/mL 2,6-DNT 2.50 mg/mL NG
6	0.200 mL of 5 mg/mL 2,6-DN7 0.250 mL of 50 mg/mL NG	10 mL	0.10 mg/ml 2,6-DNT 1.25 mg/mL NG
7	0.100 mL of 5 mg/mL 2,6-DNT 0.125 mL of 50 mg/mL NG	10 mL	0.05 mg/ml 2,6-DNT 0.62 mg/mL NG
8	0.025 mL of 5 mg/mL 2,6-DNT 0.031 mL of 50 mg/mL NG	5 mL	0.025 mg/mL 2,6-DNT 0.31 mg/mL NG

 $400~\mu\text{L}$ of spike solutions 8, 7, 6, 4, 2 spiked onto $10~\text{cm}^2$ concrete and metal surfaces

 $200~\mu L$ of spike solutions 7, 6, 5, 3, 1 spiked onto $10~\text{cm}^2$ brick and transite surfaces

Concentration on surface after spiking: NG - 125 μ g, 250 μ g, 500 μ g, 1250 μ g

2,6-DNT - 10 μg, 20 μg, 40 μg, 100 μg, 200 μg.

5. Procedure

Extraction A

- 1. Spike 10 cm^2 surface sample with acetonitrile spike solution (volume dependent on surface type). Allow solvent to evaporate.
- 2. Transfer sample to 8 ounce jar and add 20 mL $\mathrm{CH_3}\,\mathrm{CN}$. Cover jar with teflon lined cap.
- 3. Sonicate for 10 minutes.
- 4. Transfer extract A to 50 mL volumetric flask, add 25 mL H_2 0 and bring to volume with CH_3 CN.

Save surface sample for Extraction B.

Extraction B

- Add 20 mL CH₃ CN to jar with surface sample. Sonicate for 10 minutes.
- 2. Transfer surface sample to a second jar. Add 20 mL $\mathrm{CH_3}$ CN and sonicate for 10 minutes more.
- 3. Combine extracts from steps 1 and 2 and evaporate using nitrogen to less than 5 mL \cdot
- 4. Transfer evaporated extract to 10 mL volumetric flask. Add 5.0 mL $_{2}^{1}$ 0 and bring to volume with CH $_{3}^{2}$ CN. Extract B ready for analysis.

6. Calculations

Calculate found concentration for each analyte in each sample extract from daily calibration data.

Multiply found concentration by extract volume to find total μg in extract. Combine total μg in extracts A and B to find total μg on surface.

7. References

Lakings, D.B., Baker, R.J., and Crook, M.V., "Precision and Accuracy Assessment of the High Performance Liquid Chromatographic Analytical Technique for the Determination of Nitrobenzene (NB), 2,6-Dinitrotoluene (2,6-DNT), Nitroglycerin (NG), and Picric Acid (PA), Midwest Research Institute Technical Report No. 2, USATHAMA Contract No. DAAK11-81-C-0007, May, 1981.

QUANTITATIVE METHOD FOR THE DETERMINATION OF PETN ON SURFACES

1. Application

Method used to extract pentaerythrite tetranitrate (PETN) from metal, brick, concrete, transite surfaces.

A. Tested Concentration Range:

PETN 5.0 μ g/cm² to 100.0 μ g/cm²

B. Sensitivity

Instrument response for PETN is given below:

Concentration Response

0.50 µg/mL 98025 area units

C. Detection Limit

See Table II-1.

D. Interferences

There were no interferences.

E. Analysis Rate

Six samples can be extracted and prepared for analysis in three hours. Rate of analysis is given below, excluding calibration standards:

PETN 32 samples in an 8 hour day

2. Chemistry

Pentaerythrite tetranitrate C5H8N4O12 CAS RN 78-11-5 MP 140-141 C

3. Apparatus

A. Instrumentation

Beckman Model 110A Solvent Metering Pump Waters Associates Model 450 Variable Wavelength Detector Waters Associates Model U6K Injector Hewlett Packard 3390A Integrator/Recorder

B. Parameters

Column: Spherisorb ODS, 5μ , 250×4.6 mm ID Precolumn: Pellicular LC-18, 40μ , 50×4.6 mm ID

Solvent System: 65% CH₃CN/35% H₂O

Detector: UV at 230 nm Flow Rate: 1.0 mL/min Attenuation: 0.01 AUFS Injection Volume: 100 µL

C. Hardware/Glassware

Westinghouse Ultrasonic cleaner 8 ounce jars with teflon lined caps 25 mL graduated cylinders microliter syringes volumetric flasks - 50, 10, 5 mL vials - WISP and 14 mL, with teflon lined caps

D. Chemicals

Acetonitrile, HPLC grade Standard Analytical Reference Material for PETN

4. Standards

A. Calibration Standards:

Prepare stock solution as follows: 200 µL of SARM (50 mg/mL)_in 10 mL CH₃CN = 1.0 mg/mL

Calibration Standards prepared in 10 mL volumetric flasks adding 50% $\rm H_2O/50\%$ CH₃CN.

Cal. Std	ul Stock added	Concentration PETN
1	5	0.5 μg/mL
2	10	1.0 ug/mL
3	20	$2.0 \mu \text{g/mL}$
4	40	4.0 µg/mL
5	100	$10.0 \mu \text{g/mL}$
6	200	20.0 μ g /mL

B. Control Spikes

Spike solutions prepared following chart below:

Spike Solution	Amount Stock	Dilute with CH ₃ CN to	Concentration
1	12.5 µL of 50 mg/mL	5 mL	0.125 mg/mL
2	50 μL of 50 mg/mL	10 mL	0.25 mg/mL
3	100 μL of 50 mg/mL	10 mL	0.50 mg/mL
4	100 µL of 50 mg/mL	5 mL	1.0 mg/mL
5	125 µL of 50 mg/mL	5 mL	1.25 mg/mL
6	500 μL of 50 mg/mL	10 mL	2.5 mg/mL
7	500 μ L of 50 mg/mL	5 mL	5.0 mg/mL

400 μL spike solutions 1,2,3,5 and 6 spiked onto 10 cm² metal surfaces

200 μ L spike solutions 2,3,4,6,7 spiked onto 10 cm² concrete, transite, and brick surfaces.

Concentration of analytes on surface after spiking: 50 μg , 100 μg , 200 μg , 500 μg , 1000 μg .

5. Procedure

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- 1. Spike 10 cm² surface sample with acetonitrile spike solution (volume dependent on surface type). Allow solvent to evaporate.
- 2. Transfer sample to 8 ounce jar.
- 3. Add 12 mL Ch₃CN Cover jar with teflon lined cap. Sonicate for 10 min.
- 4. Transfer extract to 50 mL volumetric flask.
- Repeat steps 3 and 4 twice, adding extracts to same 50 mL volumetric flask.
- 6. Rinse jar with 17 mL ${\rm H}_2{\rm O}$ and transfer to volumetric flask.
- 7. Bring to volume with CH3CN.
- 8. Ready for HPLC analysis.

6. Calculations

Calculate found concentration for each analyte in each sample extract from daily calibration data.

Multiply found concentration by extract volume to find total ug in extract.

7. References

None

TABLE IV-4. RESULTS OF ANALYSES FOR RDX, TNB. TNT, AND TETRYL IN CONDUCTIVE NON-SPARKING FLOORING SAMPLES

	Total Micrograms			
Sample	RDX	TNB	TNT	Tetryl
Roll Flooring	48	ND	23	5.4
Roll Flooring (duplicate)	34	ND	7.3	3.2
Travelled Flooring	1200	3.6	890	ND
Travelled Flooring (duplicate)	750	15	2600	ND

ND = Not Detected

heterogeneously throughout the wood. Changes in the sample preparation procedures or analytical conditions which would circumvent this problem were not identified. In any case, further developmental effort should be based on knowledge of what wood type (i.e., species) and conditions (age, moisture content, etc.) best represent actual field conditions.

C

V. CONCLUSIONS AND RECOMMENDATIONS

- Review of the available literature suggests that available methods for the sampling and analysis of explosives/explosive residues have been developed principally for the identification of post-blast residue and for the detection of concealed bulk explosives. Most of the work in these areas has been directed toward the application of classical spot test methods, thin-layer chromatographic methods, and vapor phase detection methods. None of these methods satisfies all of the Army's requirements for the sampling and analysis of explosives/explosive residues on building materials surfaces.
- A method for the detection of explosives/explosive residues on building materials surfaces based on the formulation of charge-transfer complexes between the explosives and anthracene applied to the surface with visual identification has been developed and evaluated in the field. This method offers the following advantages:
 - (1) Sensitivity down to concentrations of analyte(s) on the order of micrograms per square centimeter of surface;
 - (2) Speed;
 - (3) Manageable hazard during and subsequent to use as compared to similar spot test methods; and
 - (4) Reversibility: the charge-transfer complexes formed in this application can be destroyed relatively easily leaving the original analyte intact and available for subsequent quantitative testing.

Field evaluations performed at two Army Ammunition Plants confirm that this method is, in fact, capable of detecting explosives contamination under field conditions, and that the presence of dirt and debris of the type under such conditions apparently does not result in false positive findings.

- Methods for the determination of explosives/explosive residues on building materials surfaces based on solvent extraction using ultrasonic agitation and analysis by high pressure liquid chromatography have been developed and evaluated in the laboratory. For the majority of analyte-surface combinations studied, analyte recoveries of 60-95% and detection limits on the order of micrograms per square centimeter of surface were obtained.
- The theoretical and practical feasibility of a method for the detection of explosives/explosive residues on building materials surfaces based on UV irradiation with subsequent detection has been demonstrated. This approach may provide the means for "scanning" of an area on a real-time basis to determine whether explosives are present. Further development of this approach is recommended.

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